Breast Cancer Progression Signatures

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FIELD OF THE INVENTION

The invention relates to the identification and use of gene expression profiles, or patterns, involved in breast cancer progression. The gene expression profiles, whether embodied in nucleic acid expression, protein expression, or other expression formats, are used in the study and/or diagnosis of cells and tissue during breast cancer progression as well as for the study and/or determination of prognosis of a patient. When used for diagnosis or prognosis, the profiles are used to predict the status and/or phenotype of cells and tissues relative to breast cancer and the treatment thereof.

BACKGROUND OF THE INVENTION

Breast cancer is by far the most common cancer among women. Each year, more than 180,000 and 1 million women in the U.S. and worldwide, respectively, are diagnosed with breast cancer. Breast cancer is the leading cause of death for women between ages 50-55, and is the most common non-preventable malignancy in women in the Western Hemisphere. An estimated 2,167,000 women in the United States are currently living with the disease (National Cancer Institute, Surveillance Epidemiology and End Results (NCI SEER) program, *Cancer Statistics* sd-71385

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Review (CSR), www-seer.ims.nci.nih.gov/Publications/CSR1973 (1998)). Based on cancer rates from 1995 through 1997, a report from the National Cancer Institute (NCI) estimates that about 1 in 8 women in the United States (approximately 12.8 percent) will develop breast cancer during her lifetime (NCI's Surveillance, Epidemiology, and End Results Program (SEER) publication SEER Cancer Statistics Review 1973-1997). Breast cancer is the second most common form of cancer, after skin cancer, among women in the United States. An estimated 250,100 new cases of breast cancer are expected to be diagnosed in the United States in 2001. Of these, 192,200 new cases of more advanced (invasive) breast cancer are expected to occur among women (an increase of 5% over last year), 46,400 new cases of early stage (in situ) breast cancer are expected to occur among women (up 9% from last year), and about 1,500 new cases of breast cancer are expected to be diagnosed in men (Cancer Facts & Figures 2001 American Cancer Society). An estimated 40,600 deaths (40,300 women, 400 men) from breast cancer are expected in 2001. Breast cancer ranks second only to lung cancer among causes of cancer deaths in women. Nearly 86% of women who are diagnosed with breast cancer are likely to still be alive five years later, though 24% of them will die of breast cancer after 10 years, and nearly half (47%) will die of breast cancer after 20 years.

Every woman is at risk for breast cancer. Over 70 percent of breast cancers occur in women who have no identifiable risk factors other than age (U.S. General Accounting Office. Breast Cancer, 1971-1991: Prevention, Treatment and Research. GAO/PEMD-92-12; 1991). Only 5 to 10% of breast cancers are linked to a family history of breast cancer (Henderson IC, Breast Cancer. In: Murphy GP, Lawrence WL, Lenhard RE (eds). *Clinical Oncology*. Atlanta, GA: American Cancer Society; 1995:198-219).

Each breast has 15 to 20 sections called lobes. Within each lobe are many smaller lobules. Lobules end in dozens of tiny bulbs that can produce milk. The lobes, lobules, and bulbs are all linked by thin tubes called ducts. These ducts lead to the nipple in the center of a dark area of skin called the areola. Fat surrounds the lobules and ducts. There are no muscles in the breast, but muscles lie under each breast and cover the ribs. Each breast also contains blood

vessels and lymph vessels. The lymph vessels carry colorless fluid called lymph, and lead to the lymph nodes. Clusters of lymph nodes are found near the breast in the axilla (under the arm), above the collarbone, and in the chest.

Breast tumors can be either benign or malignant. Benign tumors are not cancerous, they do not spread to other parts of the body, and are not a threat to life. They can usually be removed, and in most cases, do not come back. Malignant tumors are cancerous, and can invade and damage nearby tissues and organs. Malignant tumor cells may metastisize, entering the bloodstream or lymphatic system. When breast cancer cells metastisize outside the breast, they are often found in the lymph nodes under the arm (axillary lymph nodes). If the cancer has reached these nodes, it means that cancer cells may have spread to other lymph nodes or other organs, such as bones, liver, or lungs.

Major and intensive research has been focussed on early detection, treatment and prevention. This has included an emphasis on determining the presence of precancerous or cancerous ductal epithelial cells. These cells are analyzed, for example, for cell morphology, for protein markers, for nucleic acid markers, for chromosomal abnormalities, for biochemical markers, and for other characteristic changes that would signal the presence of cancerous or precancerous cells. This has led to various molecular alterations that have been reported in breast cancer, few of which have been well characterized in human clinical breast specimens. Molecular alterations include presence/absence of estrogen and progesterone steroid receptors, HER-2 expression/amplification (Mark HF, et al. HER-2/neu gene amplification in stages I-IV breast cancer detected by fluorescent in situ hybridization. Genet Med; 1(3):98-103 1999), Ki-67 (an antigen that is present in all stages of the cell cycle except G0 and used as a marker for tumor cell proliferation, and prognostic markers (including oncogenes, tumor suppressor genes, and angiogenesis markers) like p53, p27, Cathepsin D, pS2, multi-drug resistance (MDR) gene, and CD31.

Examination of cells by a trained pathologist has also been used to establish whether ductal epithelial cells are normal (i.e. not precancerous or cancerous or having another

noncancerous abnormality), precancerous (i.e. comprising hyperplasia, atypical ductal hyperplasia (ADH)) or cancerous (comprising ductal carcinoma *in situ*, or DCIS, which includes low grade ductal carcinoma *in situ*, or LG-DCIS, and high grade ductal carcinoma in situ, or HG-DCIS) or invasive (ductal) carcinoma (IDC). Pathologists may also identify the occurrence of lobular carcinoma in situ (LCIS) or invasive lobular carcinoma (ILC). Breast cancer progression may be viewed as the occurrence of abnormal cells, such as those of ADH, DCIS, IDC, LCIS, and/or ILC, among normal cells.

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It remains unclear whether normal cells become hyperplastic (such as ADH) and then progressing on to become malignant (DCIS, IDC, LCIS, and/or ILC) or whether normal cells are able to directly become malignant without transitioning through a hyperplastic stage. It has been observed, however, that the presence of ADH indicates a higher likelihood of developing a malignancy. This has resulting in treatment of patients with ADH to begin treatment with an antineoplastic/antitumor agent such as tamoxifen. This is in contrast to the treatment of patients with malignant breast cancer which usually includes surgical removal.

The rational development of preventive, diagnostic and therapeutic strategies for women at risk for breast cancer would be aided by a molecular map of the tumorigenesis process. Relatively little is known of the molecular events that mediate the transition of normal breast cells to the various stages of breast cancer progression. In particular, there is a significant paucity of information regarding the genetic changes that are associated with the earliest stages of human breast cancer, which include the transition of normal breast cells to atypical hyperplastic and/or pre-invasive malignant cells (carcinoma *in situ*).

Molecular means of identifying the differences between normal, non-cancerous cells and cancerous cells (in general) have also been the focus of intense study. The use of cDNA libraries to analyze differences in gene expression patterns in normal versus tumorigenic cells has been described (USP 4,981,783). DeRisi et al. (1996) describe the analysis of gene expression patterns between two cell lines: UACC-903, which is a tumorigenic human melanoma cell line, and UACC-903(+6), which is a chromosome 6 suppressed non-tumorigenic form of UACC-903.

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Labeled cDNA probes made from mRNA from these cell lines were applied to DNA microarrays containing 870 different cDNAs and controls. Genes that were preferentially expressed in one of the two cell lines were identified.

Golub et al. (1999) describe the use of gene expression monitoring as means to cancer class discovery and class prediction between acute myeloid leukemia (AML) and acute lymphoblastic leukemia (ALL). Their approach to class predictors used a neighborhood analysis followed by cross-validation of the validity of the predictors by withholding one sample and building a predictor based only on the remaining samples. This predictor is then used to predict the class of the withheld sample. They also used cluster analysis to identify new classes (or subtypes) within the AML and ALL.

Gene expression patterns in human breast cancers have been described by Perou et al. (1999), who studied gene expression between cultured human mammary epithelia cells (HMEC) and breast tissue samples by use of microarrays comprising about 5000 genes. They used a clustering algorithm to identify patterns of expression in HMEC and tissue samples. Perou et al. (2000) describe the use of clustered gene expression profiles to classify subtypes of human breast tumors. Hedenfalk et al. describe gene expression profiles in BRCA1 mutation positive, BRCA2 mutation positive, and sporadic tumors. Sgroi et al. also analyzed gene expression of normal and breast cancer cells from a single patient. Using gene expression patterns to distinguish breast tumor subclasses and predict clinical implications is described by Sorlie et al. and West et al.

All of the above described approaches, however, utilize heterogeneous populations of cells found in culture or in a biopsy to obtain information on gene expression patterns. The use of such populations may result in the inclusion or exclusion of multiple genes from the patterns. For this and the lack of statistical robustness reasons, the gene expression patterns observed by the above described approaches provide little confidence that the differences in gene expression may be meaningfully associated with the stages of breast cancer.



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SUMMARY OF THE INVENTION

The present invention relates to the identification and use of gene expression patterns (or profiles or "signatures") which are correlated with (and thus able to discriminate between) cells in various stages of breast cancer. Broadly defined, these stages are non-malignant versus malignant, but may also be viewed as normal versus atypical (optionally including reactive and pre-neoplastic) versus cancerous. Another definition of the stages is normal versus precancerous (e.g. atypical ductal hyperplasia (ADH) or atypical lobular hyperplasia (ALH)) versus cancerous (e.g. carcinoma *in situ* such as DCIS and/or LCIS) versus invasive (e.g. carcinomas such as IDC and/or ILC). DCIS may be further viewed as low grade versus high grade or grade I through grade III.

The gene expression patterns comprise one or more than one gene capable of discriminating between various stages of breast cancer with significant accuracy. The gene(s) are identified as correlated with various stages of breast cancer such that the levels of their expression are relevant to a determination of the stage of breast cancer of a cell. Thus in one aspect, the invention provides a method to determine the stage of breast cancer of a subject afflicted with, or suspected of having, breast cancer by assaying a cell containing sample from said subject for expression of one or more than one gene disclosed herein as correlated with one or more stages of breast cancer.

Gene expression patterns of the invention are identified by analysis of gene expression in multiple samples of each stage to be studied. The overall gene expression profile of each sample is obtained by analyzing the expressed or unexpressed state of genes in each stage relative to each other (one gene to another across all genes). This overall profile is then analyzed to identify genes that are positively, or negatively, correlated, with a stage of breast cancer relative to other genes. An expression profile of a subset of human genes may then be identified by the methods of the present invention as correlated with breast cancer. The use of multiple samples increases the confidence which which a gene may be believed to be correlated with a particular stage.

Without sufficient confidence, it remains unpredictable whether a particular gene is actually correlated with a stage of breast cancer and also unpredictable whether a particular gene may be successfully used to identify the stage of an unknown breast cancer cell sample.

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A profile of genes that are highly correlated with one stage relative to another may be used to assay an sample from a subject afflicted with, or suspected of having, breast cancer to identify the stage of breast cancer to which the sample belongs. Such an assay may be used as part of a method to determine the therapeutic treatment for said subject based upon the stage(s) of breast cancer identified.

The correlated genes may be used singly with significant accuracy or in combination to increase the ability to accurately discriminate between various stages of breast cancer. The present invention thus provides means for correlating a molecular expression phenotype with a physiological (cellular) stage or state. This correlation provides a way to molecularly diagnose and/or monitor a cell's status in comparison to different cancerous versus non-cancerous phenotypes as disclosed herein. Additional uses of the correlated gene(s) are in the classification of cells and tissues; determination of diagnosis and/or prognosis; and determination and/or alteration of therapy.

The ability to discriminate is conferred by the identification of expression of the individual genes as relevant and not by the form of the assay used to determine the actual level of expression. An assay may utilize any identifying feature of an identified individual gene as disclosed herein as long as the assay reflects, quantitatively or qualitatively, expression of the gene. Identifying features include, but are not limited to, unique nucleic acid sequences used to encode (DNA), or express (RNA), said gene or epitopes specific to, or activities of, a protein encoded by said gene. All that is required is the identity of the gene(s) necessary to discriminate between stages of breast cancer and an appropriate cell containing sample for use in an expression assay.

In one aspect, the invention provides for the identification of the gene expression patterns by analyzing global, or near global, gene expression from single cells or homogenous cell populations which have been dissected away from, or otherwise isolated or purified from, contaminating cells beyond that possible by a simple biopsy. Because the expression of numerous genes fluctuate between cells from different patients as well as between cells from the same patient sample, multiple individual gene expression patterns are used as reference data to generate models which in turn permit the identification of individual gene(s) that are most highly correlated with particular breast cancer stages and/or have the best the ability to discriminate cells of one stage from another.

In another aspect, the invention provides physical and methodological means for detecting the expression of gene(s) identified by the models generated by individual expression patterns. These means may be directed to assaying one or more aspect of the DNA template(s) underlying the expression of the gene(s), of the RNA used as an intermediate to express the gene(s), or of the proteinaceous product expressed by the gene(s).

In a further aspect, the gene(s) identified by a model as capable of discriminating between breast cancer stages may be used to identify the cellular state of an unknown sample of cell(s) from the breast. Preferably, the sample is isolated via non-invasive means. The expression of said gene(s) in said unknown sample may be determined and compared to the expression of said gene(s) in reference data of gene expression patterns from the various stages of breast cancer. Optionally, the comparison to reference samples may be by comparison to the model(s) constructed based on the reference samples.

One advantage provided by the present invention is that contaminating, non-breast cells (such as infiltrating lymphocytes or other immune system cells) are not present to possibly affect the genes identified or the subsequent analysis of gene expression to identify the status of suspected breast cancer cells. Such contamination is present where a biopsy is used to generate gene expression profiles.

While the present invention has been described mainly in the context of human breast cancer, it may be practiced in the context of breast cancer of any animal known to be potentially afflicted by breast cancer. Preferred animals for the application of the present invention are

mammals, particularly those important to agricultural applications (such as, but not limited to, cattle, sheep, horses, and other "farm animals") and for human companionship (such as, but not limited to, dogs and cats).

BRIEF DESCRIPTION OF THE FIGURES

Figure 1 is a schematic representing a data matrix of a pair-wise comparison between Grade I and Grade III DCIS among 16 samples (across the top) and a large number of genes identified by "CloneID") along the left hand side.

Figure 2 is a table showing the actual weight data corresponding to Example II, where the data from ten genes (by CloneID number vertically) are compared to DCIS and ADH samples (across the top). Some data in the table has been vertically presented to permit the table to be displayed on a single sheet. The use of "-" with data in the table reflects genes that are more highly expressed in ADH relative to DCIS. The absence of "-" reflects genes that are more highly expressed in DCIS relative to ADH.

Figure 3 is a table showing the actual weight data corresponding to Example VII, where the data from over 300 genes (by CloneID number vertically) are compared to DCIS and ADH samples (across the top). Some data in the table has been vertically presented solely for display purposes. The use of "-" with data in the table reflects genes that are more highly expressed in ADH relative to DCIS. The absence of "-" reflects genes that are more highly expressed in DCIS relative to ADH.

Figure 4 is a table showing the actual weight data corresponding to Example VIII, where the data from over 300 genes (by CloneID number vertically) are compared to samples (across the top) from two grades of DCIS. The use of "-" with data in the table reflects genes that are more

highly expressed in grade I relative to grade III. The absence of "-" reflects genes that are more highly expressed in grade III relative to grade I.

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DETAILED DESCRIPTION OF THE SPECIFIC EMBODIMENTS

Definitions of terms as used herein:

A gene expression "pattern" or "profile" or "signature" refers to the relative expression of a gene between two or more stages of breast cancer which is correlated with being able to distinguish between said stages.

A "gene" is a polynucleotide that encodes a discrete product, whether RNA or proteinaceous in nature. It is appreciated that more than one polynucleotide may be capable of encoding a discrete product. The term includes alleles and polymorphisms of a gene that encodes the same product, or a functionally associated (including gain, loss, or modulation of function) analog thereof, based upon chromosomal location and ability to recombine during normal mitosis.

A "stage" or "stages" (or equivalents thereof) of breast cancer refer to a physiologic state of a breast cell as defined by known cytological or histological (including immunohistology, histochemistry, and immunohistochemistry) procedures and are readily known to skilled in the art. Non-limiting examples include normal versus abnormal, non-cancerous versus cancerous, the different stages described herein (e.g. hyperplastic, carcinoma, and invasive), and grades within different stages (e.g. grades I, II, or III or the equivalents thereof within cancerous stages).

The terms "correlate" or "correlation" or equivalents thereof refer to an association between expression of one or more genes and a physiologic state of a breast cell to the exclusion of one or more other stages and/or identified by use of the methods as described herein. A gene may be expressed at higher or lower levels and still be correlated with one or more breast cancer stages.

A "polynucleotide" is a polymeric form of nucleotides of any length, either ribonucleotides or deoxyribonucleotides. This term refers only to the primary structure of the molecule. Thus, this term includes double- and single-stranded DNA and RNA. It also includes known types of modifications including labels known in the art, methylation, "caps", substitution of one or more of the naturally occurring nucleotides with an analog, and internucleotide modifications such as uncharged linkages (e.g., phosphorothioates, phosphorodithioates, etc.), as well as unmodified forms of the polynucleotide.

The term "amplify" is used in the broad sense to mean creating an amplification product can be made enzymatically with DNA or RNA polymerases. "Amplification," as used herein, generally refers to the process of producing multiple copies of a desired sequence, particularly those of a sample. "Multiple copies" mean at least 2 copies. A "copy" does not necessarily mean perfect sequence complementarity or identity to the template sequence.

By corresponding is meant that a nucleic acid molecule shares a substantial amount of sequence identity with another nucleic acid molecule. Substantial amount means at least 95%, usually at least 98% and more usually at least 99%, and sequence identity is determined using the BLAST algorithm, as described in Altschul et al. (1990), J. Mol. Biol. 215:403-410 (using the published default setting, i.e. parameters w=4, t=17). Methods for amplifying mRNA are generally known in the art, and include reverse transcription PCR (RT-PCR) and those described in U.S. Patent Application (number to be assigned) entitled "Nucleic Acid Amplification" filed on October 25, 2001 as attorney docket number 485772002900 as well as U.S. Provisional Patent Applications 60/298,847 (filed June 15, 2001) and 60/257,801 (filed December 22, 2000), all of which are hereby incorporated by reference in their entireties as if fully set forth. Alternatively, RNA may be directly labeled as the corresponding cDNA by methods known in the art.

A "microarray" is a linear or two-dimensional array of preferably discrete regions, each having a defined area, formed on the surface of a solid support such as, but not limited to, glass, plastic, or synthetic membrane. The density of the discrete regions on a microarray is

determined by the total numbers of immobilized polynucleotides to be detected on the surface of a single solid phase support, preferably at least about 50/cm², more preferably at least about $100/\text{cm}^2$, even more preferably at least about $500/\text{cm}^2$, but preferably below about $1,000/\text{cm}^2$. Preferably, the arrays contain less than about 500, about 1000, about 1500, about 2000, about 2500, or about 3000 immobilized polynucleotides in total. As used herein, a DNA microarray is an array of oligonucleotides or polynucleotides placed on a chip or other surfaces used to hybridize to amplified or cloned polynucleotides from a sample. Since the position of each particular group of primers in the array is known, the identities of a sample polynucleotides can be determined based on their binding to a particular position in the microarray.

Because the invention relies upon the identification of genes that are over- or under-expressed, one embodiment of the invention involves determining expression by hybridization of mRNA, or an amplified or cloned version thereof, of a sample cell to a polynucleotide that is unique to a particular gene sequence. Preferred polynucleotides of this type contain at least about 20, at least about 22, at least about 24, at least about 26, at least about 28, at least about 30, or at least about 32 consecutive basepairs of a gene sequence that is not found in other gene sequences. The term "about" as used in the previous sentence refers to an increase or decrease of 1 from the stated numerical value. Even more preferred are polynucleotides of at least about 50, at least about 100, and at least about 150 basepairs of a gene sequence that is not found in other gene sequences. The term "about" as used in the preceding sentence refers to an increase or decrease of 10% from the stated numerical value.

Alternatively, and in another embodiment of the invention, gene expression may be determined by analysis of expressed protein in a cell sample of interest by use of one or more antibodies specific for one or more epitopes of individual gene products (proteins) in said cell sample. Such antibodies are preferably labeled to permit their easy detection after binding to the gene product.

The term "label" refers to a composition capable of producing a detectable signal indicative of the presence of the labeled molecule. Suitable labels include radioisotopes,

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nucleotide chromophores, enzymes, substrates, fluorescent molecules, chemiluminescent moieties, magnetic particles, bioluminescent moieties, and the like. As such, a label is any composition detectable by spectroscopic, photochemical, biochemical, immunochemical, electrical, optical or chemical means.

The term "support" refers to conventional supports such as beads, particles, dipsticks, fibers, filters, membranes and silane or silicate supports such as glass slides.

As used herein, a "breast tissue sample" or "breast cell sample" refers to a sample of breast tissue or fluid isolated from an individual suspected of being afflicted with, or at risk of developing, breast cancer. Such samples are primary isolates (in contrast to cultured cells) and may be collected by any non-invasive means, including, but not limited to, ductal lavage, fine needle aspiration, needle biopsy, the devices and methods described in U.S. Patent 6,328,709, or any other suitable means recognized in the art. Alternatively, the "sample" may be collected by an invasive method, including, but not limited to, surgical biopsy.

"Expression" and "gene expression" include transcription and/or translation of nucleic acid material.

As used herein, the term "comprising" and its cognates are used in their inclusive sense; that is, equivalent to the term "including" and its corresponding cognates.

Conditions that "allow" an event to occur or conditions that are "suitable" for an event to occur, such as hybridization, strand extension, and the like, or "suitable" conditions are conditions that do not prevent such events from occurring. Thus, these conditions permit, enhance, facilitate, and/or are conducive to the event. Such conditions, known in the art and described herein, depend upon, for example, the nature of the nucleotide sequence, temperature, and buffer conditions. These conditions also depend on what event is desired, such as hybridization, cleavage, strand extension or transcription.

Sequence "mutation," as used herein, refers to any sequence alteration in the sequence of a gene disclosed herein interest in comparison to a reference sequence. A sequence mutation includes single nucleotide changes, or alterations of more than one nucleotide in a sequence, due

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to mechanisms such as substitution, deletion or insertion. Single nucleotide polymorphism (SNP) is also a sequence mutation as used herein. Because the present invention is based on the relative level of gene expression, mutations in non-coding regions of genes as disclosed herein may also be assayed in the practice of the invention.

"Detection" includes any means of detecting, including direct and indirect detection of gene expression and changes therein. For example, "detectably less" products may be observed directly or indirectly, and the term indicates any reduction (including the absence of detectable signal). Similarly, "detectably more" product means any increase, whether observed directly or indirectly.

Unless defined otherwise all technical and scientific terms used herein have the same meaning as commonly understood to one of ordinary skill in the art to which this invention belongs.

Specific Embodiments

The present invention relates to the identification and use of gene expression patterns (or profiles or "signatures") which discriminate between (or are correlated with) cells in various stages of breast cancer. Such patterns may be determined by the methods of the invention by use of a number of reference cell or tissue samples, such as those reviewed by a pathologist of ordinary skill in the pathology of breast cancer, which reflect various stages of breast cancer. Because the overall gene expression profile differs from person to person, cancer to cancer, and cancer cell to cancer cell, correlations between certain cell states and genes expressed or underexpressed may be made as disclosed herein to identify genes that are capable of discriminating between different breast cancer states.

The present invention may be practiced with any number of genes believed, or likely to be, differentially expressed in breast cancer cells. In Example I below, approximately 12,000 genes were used to identify hundreds of genes capable of discriminating between various stages of breast cancer as shown in Examples 2-9. The identification may be made by using expression

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profiles of various homogenous normal and breast cancer cell populations, which were isolated by microdissection, such as, but not limited to, laser capture microdissection (LCM) of 100-1000 cells. Each gene of the expression profile may be assigned weights based on its ability to discriminate between two or more stages of breast cancer (see Example I). The magnitude of each assigned weight indicates the extent of difference in expression between the two groups and is an approximation of the ability of expression of the gene to discriminate between the two groups (and thus stages). The magnitude of each assigned weight also approximates the extent of correlation between expression of individual gene(s) and particular breast cancer stages.

It should be noted that merely high levels of expression in cells from a particular stage or stages does not necessarily mean that a gene will be identified as having a high absolute weight value.

Genes with top ranking weights (in absolute terms) may be used to generate models of gene expressions that would maximally discriminate between the two groups. Alternatively, genes with top ranking weights (in absolute terms) may be used in combination with genes with lower weights without significant loss of ability to discriminate between groups. Such models may be generated by any appropriate means recognized in the art, including, but not limited to, cluster analysis, supported vector machines, neural networks or other algorithm known in the art. The models are capable of predicting the classification of a unknown sample based upon the expression of the genes used for discrimination in the models. "Leave one out" cross-validation may be used to test the performance of various models and to help identify weights (genes) that are uninformative or detrimental to the predictive ability of the models. Cross-validation may also be used to identify genes that enhance the predictive ability of the models.

The gene(s) identified as correlated with particular breast cancer stages by the above models provide the ability to focus gene expression analysis to only those genes that contribute to the ability to identify a cell as being in a particular stage of breast cancer relative to another stage or stages. The expression of other genes in a breast cancer cell would be relatively unable to provide information concerning, and thus assist in the discrimination of, different stages of

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breast cancer. For example, the alpha subunit of human topoisomerase II (identified by CloneID 825470) has been found to be useful in discriminations between normal and atypical cells (ADH and DCIS and IDC and LCIS), between normal and ADH cells compared to DCIS and IDC cells, between normal and DCIS cells, between grade I and grade III DCIS cells, and between grade I and grade III IDC cells but not between normal and ADH cells (see Examples II to IX below). Thus expression of this topoisomerase II subunit would be utilized in models to discriminate between the above listed stages but not for discerning normal from ADH cells. This type of analysis is readily incorporated into algorithms used to generate models with reference gene expression data.

As will be appreciated by those skilled in the art, the models are highly useful with even a small set of reference gene expression data and can become increasingly accurate with the inclusion of more reference data although the incremental increase in accuracy will likely diminish with each additional datum. The preparation of additional reference gene expression data using genes identified and disclosed herein for discriminating between different stages of breast cancer is routine and may be readily performed by the skilled artisan to permit the generation of models as described above to predict the status of an unknown sample based upon the expression levels of those genes.

To determine the expression levels of genes in the practice of the present invention, any method known in the art may be utilized. In one preferred embodiment of the invention, expression based on detection of RNA which hybridizes to the genes identified and disclosed herein is used. This is readily performed by any RNA detection or amplification+detection method known or recognized as equivalent in the art such as, but not limited to, reverse transcription-PCR, the methods disclosed in U.S. Patent Application (number to be assigned) entitled "Nucleic Acid Amplification" filed on October 25, 2001 as attorney docket number 485772002900 as well as U.S. Provisional Patent Applications 60/298,847 (filed June 15, 2001) and 60/257,801 (filed December 22, 2000), and methods to detect the presence, or absence, of RNA stabilizing or destabilizing sequences.

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Alternatively, expression based on detection of DNA status may be used. Detection of the DNA of an indentified gene as methylated or deleted may be used for genes that have decreased expression in correlation with a particular breast cancer stage. This may be readily performed by PCR based methods known in the art. Conversely, detection of the DNA of an indentified gene as amplified may be used for genes that have increased expression in correlation with a particular breast cancer stage. This may be readily performed by PCR based, fluorescent in situ hybridization (FISH) and chromosome in situ hybridization (CISH) methods known in the art.

Expression based on detection of a presence, increase, or decrease in protein levels or activity may also be used. Detection may be performed by any immunohistochemistry (IHC) based, blood based (especially for secreted proteins), antibody (including autoantibodies against the protein) based, ex foliate cell (from the cancer) based, mass spectroscopy based, and image (including used of labeled ligand) based method known in the art and recognized as appropriate for the detection of the protein. Antibody and image based methods are additionally useful for the localization of tumors after determination of cancer by use of cells obtained by a noninvasive procedure (such as ductal lavage or fine needle aspiration), where the source of the cancerous cells is not known. A labeled antibody or ligand may be used to localize the carcinoma(s) within a patient.

A preferred embodiment using a nucleic acid based assay to determine expression is by immobilization of one or more of the genes identified herein on a solid support, including, but not limited to, a solid substrate as an array or to beads or bead based technology as known in the art. Alternatively, solution based expression assays known in the art may also be used. The immobilized gene(s) may be in the form of polynucleotides that are unique or otherwise specific to the gene(s) such that the polynucleotide would be capable of hybridizing to a DNA or RNA corresponding to the gene(s). These polynucleotides may be the full length of the gene(s) or be short sequences of the genes that are optionally minimally interrupted (such as by mismatches or

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inserted non-complementary basespairs) such that hybridization with a DNA or RNA corresponding to the gene(s) is not affected.

The immobilized gene(s) may be used to determine the state of nucleic acid samples prepared from sample breast cell(s) for which the pre-cancer or cancer status is not known or for confirmation of a status that is already assigned to the sample breast cell(s). Without limiting the invention, such a cell may be from a patient suspected of being afflicted with, or at risk of developing, breast cancer. The immobilized polynucleotide(s) need only be sufficient to specifically hybridize to the corresponding nucleic acid molecules derived from the sample. While even a single correlated gene sequence may to able to provide adequate accuracy in discriminating between two breast cancer cell stages, two or more, three or more, four or more, five or more, six or more, seven or more, eight or more, nine or more, ten or more, or eleven or more of the genes identified herein may be used as a subset capable of discriminating may be used in combination to increase the accuracy of the method. The invention specifically contemplates the selection of more than one, two or more, three or more, four or more, five or more, six or more, seven or more, eight or more, nine or more, ten or more, or eleven or more of the genes disclosed in the tables and figures herein for use as a subset in the identification of whether an unknown or suspicious breast cancer sample is normal or is in one or more stages of breast cancer. Optionally, the genes used will not include CloneID 809507, which is also known as GenBank accession number AA454563, described as an EST with high similarity to CD63 but of unknown function.

In embodiments where only one or a few genes are to be analyzed, the nucleic acid derived from the sample breast cancer cell(s) may be preferentially amplified by use of appropriate primers such that only the genes to be analyzed are amplified to reduce contaminating background signals from other genes expressed in the breast cell. Alternatively, and where multiple genes are to be analyzed or where very few cells (or one cell) is used, the nucleic acid from the sample may be globally amplified before hybridization to the immobilized

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polynucleotides. Of course RNA, or the cDNA counterpart thereof may be directly labeled and used, without amplification, by methods known in the art.

The above assay embodiments may be used in a number of different ways to identify or detect the breast cancer stage, if any, of a breast cancer cell sample from a patient. In many cases, this would reflect a secondary screen for the patient, who may have already undergone mammography or physical exam as a primary screen. If positive, the subsequent needle biopsy, ductal lavage, fine needle aspiration, or other analogous methods may provide the sample for use in the above assay embodiments. The present invention is particularly useful in combination with non-invasive protocols, such as ductal lavage or fine needle aspiration, to prepare a breast cell sample. The current analysis of ductal lavage samples is by cytological examination by a trained pathologist who classifies the samples in terms that are at least partly subjective: unsatisfactory (too few cells), benign (including fibrocystic change), atypical (or mild atypia), suspicious (or marked atypia), or malignant.

The present invention provides a more objective set of criteria, in the form of gene expression profiles of a discrete set of genes, to discriminate (or delineate) between meaningful stages (or classes) of breast cancer cells. In particularly preferred embodiments of the invention, the assays are used to discriminate between ADH and DCIS or otherwise malignant cells, which is a critical determination for decisions concerning subsequent treatment and therapy for the patient. Another particularly preferred determination is between the three grades (I, II, III) of carcinomas *in situ* as well as the discrimination between grade III carcinomas *in situ* and invasive carcinomas. Other pairwise comparisons that are provided by the invention include, but are not limited to, normal versus ADH, normal versus carcinoma *in situ*, normal versus invasive, normal versus cancerous (i.e. carcinoma present), ADH versus cancerous, and carcinoma *in situ* versus invasive. With the use of alternative algorithms, such as neural networks, comparisons that discriminate between multiple (more than pairwise) classes may also be performed. It is believed by the inventors that the present invention is the first example of objective, molecular criteria for making these discriminations.

In one embodiment of the invention, the isolation and analysis of a breast cancer cell sample may be performed as follows:

- (1) Ductal lavage or other non-invasive procedure is performed on a patient to obtain a sample.
- (2) Sample is prepared and coated onto a microscope slide. Note that ductal lavage results in clusters of cells that are cytologically examined as stated above.
- (3) Pathologist or image analysis software scans the sample for the presence of atypical cells.
- (4) If atypical cells are observed, those cells are harvested (e.g. by microdissection such as LCM).
- (5) RNA is extracted from the harvested cells.
- (6) RNA is purified, amplified, and labeled.
- (7) Labeled nucleic acid is contacted with a microarray containing polynucleotides of the genes identified herein as correlated to discriminations between two or more stages of breast cancer under hybridization conditions, then processed and scanned to obtain a pattern of intensities of each spot (relative to a control for general gene expression in cells) which determine the level of expression of the gene(s) in the cells.
- (8) The pattern of intensities is analyzed by comparison to the expression patterns of the genes in known samples of normal and breast cancer cells (relative to the same control).

A specific example of the above method would be performing ductal lavage following a primary screen, observing and collecting atypical cells for analysis. The comparison to known expression patterns, such as that made possible by a model generated by an algorithm (such as, but not limited to nearest neighbor type analysis, SVM, or neural networks) with reference gene expression data for the different breast cancer stages, identifies the cells as being most likely ADH.

Alternatively, the sample may permit the collection of both normal and atypical cells for analysis. The gene expression patterns for each of these two samples will be compared to each sd-71385

other as well as the model and the normal versus individual abnormal comparisons therein based upon the reference data set. This approach can be significantly more powerful that the atypical cells only approach because it utilizes significantly more information from the normal cells and the differences between normal and atypical cells (in both the sample and reference data sets) to determine the status of the atypical cells from the sample.

By appropriate selection of the genes used in the analysis, identification of the relative amounts of atypical cells may also be possible, although in most clinical settings, the identification of the highest grade of breast cancer with confidence makes identification of lower grades less important. Stated differently, the identification of invasive cancer determines the clinical situation regardless of the presence of carcinoma *in situ* or hyperplastic cells, or the identification of carcinoma *in situ* makes determines the clinical situation regardless of the presence of hyperplastic cells.

With use of the present invention, skilled physicians may prescribe treatments based on non-invasive samples that they would have prescribed for a patient which had previously received a diagnosis via a solid tissue biopsy.

The above discussion is also applicable where a palpable lesion is detected followed by fine needle aspiration or needle biopsy of cells from the breast. The cells are plated and reviewed by a pathologist or automated imaging system which selects cells for analysis as described above. This again provides a means of linking visual to molecular cytology and provides a less subjective means of identifying the physiological state of breast cancer cells without the need for invasive solid tissue biopsies.

The present invention may also be used, however, with solid tissue biopsies. For example, a solid biopsy may be collected and prepared for visualization followed by determination of expression of one or more genes identified herein to determine the stage of breast cancer, if any. One preferred means is by use of *in situ* hybridization with polynucleotide or protein identifying probe(s) for assaying expression of said gene(s).

In an alternative method, the solid tissue biopsy may be used to extract molecules followed by analysis for expression of one or more gene(s). This provides the possibility of leaving out the need for visualization and collection of only those cells suspected of being atypical. This method may of course be modified such that only cells suspected of being atypical are collected and used to extract molecules for analysis. This would require visualization and selection as an prerequisite to gene expression analysis.

In a further modification of the above, both normal cells and cells suspected of being atypical are collected and used to extract molecules for analysis of gene expression. The approach, benefits and results are as described above using non-invasive sampling.

In a further alternative to all of the above, the gene(s) identified herein may be used as part of a simple PCR or array based assay simply to determine the presence of atypical cells in a sample from a non-invasive sampling procedure. This is simple to perform and utilizes genes identified to be the best discriminators of normal versus abnormal cells without the need for any cytological examination. If no atypical cells are identified, no cytological examination is necessary. If atypical cells are identified, cytological examination follows, and a more comprehensive analysis, as described above, may follow.

The genes identified herein may be used to generate a model capable of predicting the breast cancer stage (if any) of an unknown breast cell sample based on the expression of the identified genes in the sample. Such a model may be generated by any of the algorithms described herein or otherwise known in the art as well as those recognized as equivalent in the art using gene(s) (and subsets thereof) disclosed herein for the identification of whether an unknown or suspicious breast cancer sample is normal or is in one or more stages of breast cancer. The model provides a means for comparing expression profiles of gene(s) of the subset from the sample against the profiles of reference data used to build the model. The model can compare the sample profile against each of the reference profiles or against model defining delineations made based upon the reference profiles. Additionally, relative values from the sample profile may be used in comparison with the model or reference profiles.

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In a preferred embodiment of the invention, breast cell samples identified as normal and abnormal (atypical) from the same subject may be analyzed for their expression profiles of the genes used to generate the model. This provides an advantageous means of identifying the stage of the abnormal sample based on relative differences from the expression profile of the normal sample. These differences can then be used in comparison to differences between normal and individual abnormal reference data which was also used to generate the model.

The detection of gene expression from the samples may be by use of a single microarray able to assay gene expression from all pairwise comparisons disclosed herein for convenience and accuracy.

Other uses of the present invention include providing the ability to identify breast cancer cell samples as being those of a particular stage of cancer for further research or study. This provides a particular advantage in many contexts requiring the identification of breast cancer stage based on objective genetic or molecular criteria rather than cytological observation. It is of particular utility to distinguish different grades of a particular breast cancer stage for further study, research or characterization because no objective criteria for such delineation was previously available.

The materials for use in the methods of the present invention are ideally suited for preparation of kits produced in accordance with well known procedures. The invention thus provides kits comprising agents for the detection of expression of the disclosed genes for identifying breast cancer stage. Such kits optionally comprising the agent with an identifying description or label or instructions relating to their use in the methods of the present invention, is provided. Such a kit may comprise containers, each with one or more of the various reagents (typically in concentrated form) utilized in the methods, including, for example, pre-fabricated microarrays, buffers, the appropriate nucleotide triphosphates (e.g., dATP, dCTP, dGTP and dTTP; or rATP, rCTP, rGTP and UTP), reverse transcriptase, DNA polymerase, RNA polymerase, and one or more primer complexes of the present invention (e.g., appropriate length

poly(T) or random primers linked to a promoter reactive with the RNA polymerase). A set of instructions will also typically be included.

The methods provided by the present invention may also be automated in whole or in part. All aspects of the present invention may also be practiced such that they consist essentially of a subset of the disclosed genes to the exclusion of material irrelevant to the identification of breast cancer stages in a cell containing sample.

Having now generally described the invention, the same will be more readily understood through reference to the following examples which are provided by way of illustration, and are not intended to be limiting of the present invention, unless specified.

EXAMPLES

Example I: Materials and Methods

Clinical specimens

Clinical biopsies from 30 patients were obtained from the Massachusetts General Hospital with Institutional Review Board approval. The tissue from one of the patients was not associated with breast cancer of any kind since it was from a breast reduction procedure. Pathological and histological information for the biopsies were also obtained. Three independent captures of about 1000 breast epithelial cells of one or more of the four different disease stages (normal, N; atypical ductal hyperplasia, ADH; ductal carcinoma *in situ*, DCIS; invasive ductal carcinoma, IDC) were procured from each biopsy using Laser Capture Microdissection (LCM, Arcturus Engineering). Three independent captures of LCIS (lobular carcinoma in situ) in one biopsy were also made. Total RNA was extracted from the captured (procured) cells and amplified with a T7-promoter based RNA amplification protocol. The human universal

reference RNA (Stratagene, La Jolla), was similarly amplified and used as the reference channel in a two-color microarray hybridization.

Microarrays

To maximize coverage of breast cancer-related genes on the microarrays used, 11,435 cDNA clones from the IMAGE consortium (Research Genetics) were obtained. These clones were selected based on literature knowledge (such as, but not limited to, preferential expression in cancer versus normal cells) and after mining (such as, but not limited to, preferential expression in breast tissues) gene expression information in the expressed sequence tags (EST) databases and the Serial Analysis of Gene Expression (SAGE) data sets available from the National Center for Biotechnology Information (NCBI, http://www/ncbi.nlm.nih.gov).

Microarray data processing

Microarray images were analyzed with ImaGene (BioDiscovery) to find and quantitate each spot on the microarray. Spots flagged by ImaGene as poor spots using standard criteria used with the software for the standardization of signals were excluded from further analysis. Raw Cy5 (sample channel) and Cy3 (reference channel) intensities and associated local background estimates for each spot were then examined. The signal/noise ratio, defined as the spot intensity over background intensity, was used as the second criteria for spot exclusion; spots with signal/noise ratio < 3.0 in the reference channel or < 1.5 in the sample channel were excluded from further analysis. Background-subtracted intensities across the chip were normalized to the 75th-percentile of the spot intensity distribution on the entire chip (alternative normalizations to the mean, median or other point may also be used as known in the art). Cy5/Cy3 ratios of each spot for each cellular state were averaged across each of six measurements (3 LCM captures x 2 chips/capture = 6 chips); outliers among the 6 data points were removed before taking the average. The resulting data were formatted as a data matrix

(samples along the top horizontal axis and gene identity along the vertical axis) for data mining (see Figure 1 with data).

Microarray data analysis

Before further analysis, each value in a row (gene) of the gene expression matrix was divided by the median value for the row, and the resulting matrix log-transformed. Normalized, median-centered, and log-transformed, gene expression data matrix was loaded into GeneMaths software (Applied-Maths, Belgium). Clustering and discriminant analysis were performed to identify sets of genes associated with different cellular states. For each pair-wise comparison between two breast cancer stages, samples are assigned to either the positive group or negative group, and genes were sorted by their discriminatory weights. The absolute value of the weight of a gene indicates the extent of difference in expression between the two groups; the positively signed genes are expressed higher in one group and the negatively singed genes are expressed higher in the other group.

The utility of the top-ranking genes as a diagnostic test was evaluated using the support vector machines (SVMs) algorithm (see Yeang, C. H., S. Ramaswamy, et al. (2001). "Molecular classification of multiple tumor types." Bioinformatics 17 Suppl 1: S316-22; Xiong, M., X. Fang, et al. (2001). "Biomarker identification by feature wrappers." Genome Res 11(11): 1878-87this one used linear discriminate analysis, logistic regression and svm; Furey, T. S., N. Cristianini, et al. (2000). "Support vector machine classification and validation of cancer tissue samples using microarray expression data." Bioinformatics 16(10): 906-14; and Brown, M. P., W. N. Grundy, et al. (2000). "Knowledge-based analysis of microarray gene expression data by using support vector machines." Proc Natl Acad Sci U S A 97(1): 262-7, who state "SVMs are considered a supervised computer learning method because they exploit prior knowledge of gene function to identify unknown genes of similar function from expression data. SVMs avoid several problems associated with unsupervised clustering methods, such as hierarchical

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clustering and self-organizing maps.") Other algorithms, such as, but not limited to, linear discriminate analysis, logistic regression, cluster analysis, K-th nearest neighbor, or neural nets.

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The support vector machines algorithm finds the maximal margin hyperplane that separate the two groups under comparison. The method of leave-one-out cross-validation was used to test the performance of a given set of genes; one sample was taken in turn out of the training set and a model is built using the rest of the training set, which is in then applied to classify the left-out sample. The accuracy of the genes in the cross-validation procedure is the percentage of correct classifications over the total number of the training samples.

Example II: 10 Genes for discriminating between ADH and DCIS

Based upon the methodology of Example I above, 10 genes identified as capable of discriminating between ADH and DCIS are listed in Table 1 below along with a brief description of the gene. CloneID as used in the context of the present invention refers to the IMAGE Consortium clone ID number of each gene, the sequences of which are hereby incorporated by reference in their entireties as they are available from the Consortium at http://image.llnl.gov/ as accessed on the filing date of the present application. Weight refers to the absolute value indicating the extent of difference in expression between ADH and DCIS where the positively signed values are expressed higher in ADH and the negatively signed values are expressed higher in DCIS; Chromosome Location refers to the human chromosome to which the gene has been assigned, and Description provides a brief identifier of what the gene encodes. The actual data corresponding to the assigned weights are shown in Figure 2.

Table 1

GenelD	Weight	Chromosome Location	Description
825470	0.9946555	17q21-q22	topoisomerase (DNA) II alpha (170kD) hypothetical protein baculoviral IAP repeat-containing 5
595213	0.9884884	8	
796694	0.9852686	17q25	(survivin) parathyroid hormone-like hormone SPARC-like 1 (mast9, hevin) peanut (Drosophila)-like 2 frizzled-related protein ESTs, Weakly similar to 2004399A
1404774	-1.5638738	12p12.1-p11.2	
823871	-1.5260464	7	
1882697	-1.401878	17q22-q23	
140071	-1.3881954	2	
160192	-1.2551663	5	chromosomal protein [H.sapiens] ets variant gene 5 (ets-related molecule) troponin I, skeletal, fast
796542	-1.1401853	3q28	
611532	-1.1227597	11p15.5	

Example III: Genes for discriminating between normal and non-normal (a combination of ADH, DCIS, and IDC) cells from breast tissue

As shown in Table 2 below, 850 genes were identified as being able to discriminate between normal and "abnormal" (defined in this instance as any sample that was not normal), which includes ADH, DCIS and IDC.

Table 2

ClonelD	Weight	Description
1323448	1.5470535	cysteine-rich protein 1 (intestinal)
788654	1.4818381	growth factor receptor-bound protein 2
35147	1.3764654	"ESTs, Weakly similar to unnamed protein product
		[H.sapiens]"
745606	1.3470375	hypothetical protein PP591
1500000	1.3204029	"H2B histone family, member B"
595037	1.3057353	retinoic acid induced 3

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565319	1.2369212	Homo sapiens mRNA; cDNA DKFZp564B1264 (from clone DKFZp564B1264)
122077 1609836		putative membrane protein glutamate-ammonia ligase (glutamine synthase)
1505038 178805		hypothetical protein FLJ20171 "Human DNA sequence from clone RP5-850E9 on chromosome 20. Contains part of the gene for a novel C2H2 type zinc finger protein similar to Drosophila Scratch (Scrt), Slug and Xenopus Snail, a novel gene similar to Drosophila CG6762, STSs, GSSs and five CpG
812238 1492238 366132	1.1991653	hypothetical protein MGC4692 HSPC003 protein "succinate dehydrogenase complex, subunit C, integral membrane protein, 15kD"
796469	1.1850928	HSPC150 protein similar to ubiquitin-conjugating enzyme
488964 471568		"H2A histone family, member O" hematological and neurological expressed 1
1554549 283919 1492463 199403	1.118257	hydroxyacyl glutathione hydrolase "H2A histone family, member L" "selenoprotein X, 1" "lectin, galactoside-binding, soluble, 8 (galectin 8)"
796723	1.0724133	Homo sapiens clone CDABP0014 mRNA sequence
138189 1574058		Wolfram syndrome 1 (wolframin) "1-acylglycerol-3-phosphate O-acyltransferase 2 (lysophosphatidic acid acyltransferase, beta)"
811774 359887		CGI-49 protein translocase of inner mitochondrial membrane 17 (yeast) homolog A
1709791 244801 1917941 1858892 288999	1.0162794 0.9960315 0.9897362	BAI1-associated protein 1 Rho guanine exchange factor (GEF) 11 purine-rich element binding protein B hypothetical protein MGC4825 small protein effector 1 of Cdc42

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503215 1911343 2029173	0.967716	pilin-like transcription factor "RAB26, member RAS oncogene family" "ESTs, Weakly similar to N-WASP [H.sapiens]"
470061 2016908		seven in absentia (Drosophila) homolog 2 "ESTs, Weakly similar to CA13_HUMAN COLLAGEN ALPHA 1(III) CHAIN PRECURSOR [H.sapiens]"
1469425 814054 814528	0.958989	SRY (sex determining region Y)-box 22 KIAA0040 gene product "Homo sapiens cDNA: FLJ22139 fis, clone HEP20959"
68636 290841 236034	0.9562201	hypothetical protein MGC2477 "H2B histone family, member A" "uncoupling protein 2 (mitochondrial, proton carrier)"
782428 131094		KIAA0250 gene product "Homo sapiens cDNA: FLJ21587 fis, clone COL06946"
488202	0.9501137	"ESTs, Weakly similar to YZ28_HUMAN HYPOTHETICAL PROTEIN ZAP128 [H.sapiens]"
1435862	0.9500093	"antigen identified by monoclonal antibodies 12E7, F21 and O13"
823598	0.9452282	"proteasome (prosome, macropain) 26S subunit, non- ATPase, 12"
1492426 795185		chromosome 19 open reading frame 3 xenotropic and polytropic retrovirus receptor
810558	0.9227709	"proteasome (prosome, macropain) 26S subunit, ATPase, 4"
469686	0.9203022	"Ric (Drosophila)-like, expressed in many tissues"
280375		PRO2000 protein
729975	0.9149894	meningioma expressed antigen 5 (hyaluronidase)
810124	0.9124035	"platelet-activating factor acetylhydrolase, isoform lb, gamma subunit (29kD)"
813281		WW domain-containing protein 1
39884	0.9091807	IMP (inosine monophosphate) dehydrogenase 1
76605 1636092		nesca protein hypothetical protein FLJ20657
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741474 1605426 788654		glucose phosphate isomerase hypothetical protein FLJ13352
868128		JM4 protein
308466		GTP-binding protein Sara
44292	0.8732235	Homo sapiens mRNA; cDNA DKFZp434C107 (from clone DKFZp434C107)
826256	0.8700831	transmembrane 7 superfamily member 1 (upregulated in kidney)
685516	0.8694652	putative G protein-coupled receptor
810711	0.869424	stearoyl-CoA desaturase (delta-9-desaturase)
753299		hypothetical protein FLJ10504
731044		glutaredoxin 2
824052		chromosome 6 open reading frame 1
843195	0.86501	phosphoserine phosphatase
292770	0.0070990	"Homo sapiens, clone IMAGE:3627860, mRNA, partial cds"
859761	0.8563964	poliovirus receptor-related 2 (herpesvirus entry mediator B)
347373	0.8555396	"transcription elongation factor B (SIII), polypeptide 1 (15kD, elongin C)"
1640821	0.8544081	"ESTs, Weakly similar to 178885 serine/threonine-specific protein kinase [H.sapiens]"
704414	0.8520508	small nuclear ribonucleoprotein polypeptides B and B1
810725	0.8510425	"ATPase, H+ transporting, lysosomal (vacuolar proton pump) 21kD"
782608	0.8507815	mitochondrial ribosomal protein L9
2019223		mitochondrial ribosomal protein L17
2110511	0.8471736	artemin
1474955	0.8460414	"TATA box binding protein (TBP)-associated factor, RNA polymerase II, N, 68kD (RNA-binding protein 56)"
810612	0.842284	S100 calcium-binding protein A11 (calgizzarin)
744417	0.8409538	carnitine acetyltransferase
969877		"synaptosomal-associated protein, 25kD"
272529	0.8371698	phosphomannomutase 2
768570		hypothetical protein FLJ11280
824879	0.8366695	hypothetical protein MGC11275
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2014034	0.8354357	"methylene tetrahydrofolate dehydrogenase (NAD+ dependent), methenyltetrahydrofolate cyclohydrolase"
2054635	0.831532	"proteasome (prosome, macropain) subunit, alpha type, 7"
2052113 742595 2016648	0.8291965	hypothetical protein FLJ10903 cyclin-dependent kinase 5 Homo sapiens mRNA; cDNA DKFZp434N1728 (from clone DKFZp434N1728)
1573251	0.823025	peroxisomal long-chain acyl-coA thioesterase
1869201 37708 241348 810063	0.8194577 0.8182422	hypothetical protein MGC2745 hypothetical protein MGC3101 prenylcysteine lyase "growth factor, erv1 (S. cerevisiae)-like (augmenter of liver regeneration)"
731308		citrate synthase
209066 590774	0.8120797	mitogen-activated protein kinase 13
1435003		tumor suppressing subtransferable candidate 1
502774 488505 109863 813419	0.808712 0.8066906	hypothetical protein FLJ20623 accessory proteins BAP31/BAP29 epithelial membrane protein 2 "hydroxyacyl-Coenzyme A dehydrogenase, type II"
1845169 742707		"RAB35, member RAS oncogene family" "ESTs, Weakly similar to MUC2_HUMAN MUCIN 2 PRECURSOR [H.sapiens]"
594500	0.7935958	
1456701 271472		B-cell CLL/lymphoma 9 C3HC4-like zinc finger protein
1473922		"actin related protein 2/3 complex, subunit 3 (21 kD)"
51773 898032 340558	0.7852733	hypothetical protein MGC3077 KIAA0097 gene product "actin related protein 2/3 complex, subunit 5 (16 kD)"
773922 768064		KIAA0005 gene product "cytochrome P450, subfamily I (aromatic compound-inducible), polypeptide 1"

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625923	0.7725056	phosphoenolpyruvate carboxykinase 2 (mitochondrial)
150003 741977		hypothetical protein FLJ13187 "B-factor, properdin"
365738 814350	0.7709456	
2108077		insulin-degrading enzyme CGI-112 protein
209066		serine/threonine kinase 15
564492		mitochondrial carrier homolog 2
366353		DKFZP564C186 protein
813751		"sialyltransferase 4C (beta-galactosidase alpha-2,3-sialytransferase)"
781097	0.7633668	·
233349		hypothetical protein FLJ10761
825327		"Homo sapiens cDNA FLJ14105 fis, clone
020021	0.7022414	MAMMA1001202"
248649	0.761865	
589232		hypothetical protein FLJ11506
564847	0.757516	
754628	0.7573763	
1500162	0.7571399	ESTs
1738208	0.7565056	"a disintegrin-like and metalloprotease (reprolysin type) with thrombospondin type 1 motif, 4"
842994	0.7563101	cathepsin Z
42408		hypothetical protein MGC4604
470124		RAD1 (S. pombe) homolog
809357		Bernardinelli-Seip congenital lipodystrophy 2 (seipin)
624667	0.7524325	CGI-92 protein
49351	0.75115	SEX gene
46248	0.7510436	ADP-ribosyltransferase (NAD+; poly (ADP-ribose) polymerase)
686552	0.7506994	golgi phosphoprotein 1
1631735		"Homo sapiens, clone IMAGE:3604336, mRNA, partial cds"
1536006	0.7499887	ESTs
83363	0.748646	protein-L-isoaspartate (D-aspartate) O-methyltransferase
713782	0.7484589	a disintegrin and metalloproteinase domain 15 (metargidin)

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727078	0.7472345	"Homo sapiens cDNA: FLJ23602 fis, clone LNG15735"
503889 284261		hypothetical protein FLJ10656 uncharacterized hematopoietic stem/progenitor cells protein MDS030
1473289	0.7441483	protective protein for beta-galactosidase (galactosialidosis)
49273	0.742123	"solute carrier family 27 (fatty acid transporter), member 4"
785701 41569 810402 1456348	0.7412357 0.7407907	"RAB31, member RAS oncogene family" hypothetical protein FLJ12650 hypothetical protein N-acetylneuraminic acid phosphate synthase; sialic acid
256619 743589		synthase hydroxysteroid (17-beta) dehydrogenase 7 "ESTs, Weakly similar to T2D3_HUMAN TRANSCRIPTION INITIATION FACTOR TFIID 135 KDA SUBUNIT [H.sapiens]"
1409509 122241		"troponin T1, skeletal, slow" "proteasome (prosome, macropain) subunit, beta type, 2"
470099	0.7343404	HT002 protein; hypertension-related calcium-regulated
725454 796694		gene CDC28 protein kinase 2 baculoviral IAP repeat-containing 5 (survivin)
186768	0.7298378	"Homo sapiens, clone IMAGE:3604869, mRNA"
813629 842980		YME1 (S.cerevisiae)-like 1 developmentally regulated GTP-binding protein 1
769921 2322367 951233	0.7249854	ubiquitin carrier protein E2-C reticulon 4 "proteasome (prosome, macropain) subunit, beta type, 3"
842825 782608 2043167 827171 120271 1518890	0.7230962 0.7224903 0.7211896 0.7208543	G1 to S phase transition 1 BCL2-associated athanogene 3 ESTs hypothetical protein MGC4692 "metallothionein-like 5, testis-specific (tesmin)"
60565 sd-71385	0.720547	lethal giant larvae (Drosophila) homolog 2

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2050827	0.7156518	"proteasome (prosome, macropain) 26S subunit, ATPase, 5"
1635681	0.7152705	"NADH dehydrogenase (ubiquinone) 1 alpha subcomplex, 2 (8kD, B8)"
731023	0.7151176	WD repeat domain 5
1518402	0.7150061	KIAA1361 protein
752631	0.7139862	"fibroblast growth factor receptor 3 (achondroplasia, thanatophoric dwarfism)"
785616	0.7131593	"signal sequence receptor, alpha (translocon-associated protein alpha)"
686172	0.7131363	KIAA0008 gene product
40173	0.7124913	KIAA0807 protein
123441	0.7124698	ribosomal protein L7a
123614	0.7117834	hypothetical protein MGC4675
811024		bone marrow stromal cell antigen 2
2302099	0.7117055	sialidase 3 (membrane sialidase)
756442		P450 (cytochrome) oxidoreductase
811585		huntingtin (Huntington disease)
279970		adenosine A2a receptor
1517171		"interleukin 2 receptor, alpha"
838366	0.702617	3-hydroxymethyl-3-methylglutaryl-Coenzyme A lyase (hydroxymethylglutaricaciduria)
809944	0.7021108	KIAA0310 gene product
1393018		"general transcription factor IIIC, polypeptide 1 (alpha subunit, 220kD)"
725978	0.7010756	"ESTs, Moderately similar to ALU7_HUMAN ALU SUBFAMILY SQ SEQUENCE CONTAMINATION WARNING ENTRY [H.sapiens]"
325606	0.7008219	hypothetical protein MGC14353
795256		NPD007 protein
365358	0.696963	pM5 protein
67765	0.6967324	carboxypeptidase M
358267	0.6962436	"EST, Moderately similar to AF119917 63 PRO2831 [H.sapiens]"
149355	0.6943923	translocating chain-associating membrane protein
212542	0.6938079	"Homo sapiens cDNA FLJ12900 fis, clone NT2RP2004321"
79520	0.6919579	"RAB2, member RAS oncogene family"

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739126	0.6881663	tissue specific transplantation antigen P35B
202514	0.6872212	DNA (cytosine-5-)-methyltransferase 3 alpha
1884404 814378 2018337	0.6857148	KIAA0285 gene product "serine protease inhibitor, Kunitz type, 2" "glucosidase, beta; acid (includes glucosylceramidase)"
701751	0.6824676	cut (Drosophila)-like 1 (CCAAT displacement protein)
2018084 781019 2244196 124447 789012 490778	0.6819736 0.6807351 0.679866 0.6794432	Ste-20 related kinase paraoxonase 2 B-cell receptor-associated protein BAP29 KIAA1184 protein fibulin 2 low molecular mass ubiquinone-binding protein (9.5kD)
2011515 80764 841679	0.6765616	DKFZP586B0923 protein hypothetical protein calcium and integrin binding protein (DNA-dependent protein kinase interacting protein)
108425 878406 487733 625693 1325816		metaxin 1 hypothetical protein MGC10911 polymerase (RNA) II (DNA directed) polypeptide L (7.6kD)
278531 744374 203003	0.6732551	cytochrome c oxidase subunit VIc putative ankyrin-repeat containing protein "non-metastatic cells 4, protein expressed in"
564981 812994 172517 205049	0.6701228	ESTs "retinoid X receptor, alpha" hippocalcin-like 1 protein kinase H11; small stress protein-like protein HSP22
1734309 1631699 2028949	0.6665952	sperm associated antigen 4 valosin-containing protein hypothetical protein PRO1855

530197	0.6659784	"ESTs, Moderately similar to ALU8_HUMAN ALU SUBFAMILY SX SEQUENCE CONTAMINATION WARNING ENTRY [H.sapiens]"	
199645 704254	0.6654138 0.6653635	nicastrin a disintegrin and metalloproteinase domain 8	
1592715	0.6637095	"Homer, neuronal immediate early gene, 3"	
302031 239568 770785 785795 810734 1696757 811761 150314 489351 593431	0.661735 0.6613519 0.660789 0.6603412 0.6591801 0.6587469 0.6584621 0.65846	"polymerase (DNA-directed), delta 4" hypothetical protein KIAA1165 Nijmegen breakage syndrome 1 (nibrin) lysophospholipase I	
37554 126851 265103	0.6559601	hypothetical protein FLJ22353 hypothetical protein FLJ11160 Homo sapiens mRNA; cDNA DKFZp547M123 (from clone DKFZp547M123)	
51083	0.6544472	"catenin (cadherin-associated protein), delta 2 (neural plakophilin-related arm-repeat protein)"	
431505	0.6534315	"ESTs, Highly similar to A31026 probable membrane receptor protein [H.sapiens]"	
1420370	0.6531171	biliverdin reductase B (flavin reductase (NADPH))	
1476053	0.6522237	RAD51 (S. cerevisiae) homolog (E coli RecA homolog)	
882484	0.6521026	"chaperonin containing TCP1, subunit 7 (eta)"	
51532	0.6504769	ADP-ribosylation factor-like 6 interacting protein	
280249 138788 1492468	0.6497728	Kruppel-like factor 7 (ubiquitous) prolactin receptor DEME-6 protein	

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221295	0.6477514	"inhibitor of DNA binding 2, dominant negative helix-loophelix protein"
897806	0.6470379	"hypoxia-inducible factor 1, alpha subunit (basic helix-loophelix transcription factor)"
358936	0.6441198	"ESTs, Weakly similar to T2D3_HUMAN TRANSCRIPTION INITIATION FACTOR TFIID 135 KDA SUBUNIT
840865	0.644011	[H.sapiens]" "myristoylated alanine-rich protein kinase C substrate (MARCKS, 80K-L)"
788334 366067		mitochondrial ribosomal protein L23 cerebellar degeneration-related protein (62kD)
825585 1460110		tubulin-specific chaperone e "proteasome (prosome, macropain) subunit, beta type, 5"
344091 196189 2248488		ESTs cytochrome b-5 ems1 sequence (mammary tumor and squamous cell carcinoma-associated (p80/85 src substrate)
509823	0.6334191	carcinoembryonic antigen-related cell adhesion molecule 6 (non-specific cross reacting antigen)
144880	0.6332999	hypothetical protein from EUROIMAGE 1759349
755599	0.6330221	interferon induced transmembrane protein 1 (9-27)
207288 1639531 144740	0.6315732	insulin induced gene 1 "RAB27A, member RAS oncogene family" similar to phosphatidylcholine transfer protein 2
814306 301122 124298 1733262 376875	0.6299499 0.6294358 0.6293971	tumor protein D52 extracellular matrix protein 1 microsomal glutathione S-transferase 3 BLu protein flavin containing monooxygenase 1
773188	0.6270494	"nuclear receptor subfamily 1, group D, member 2"
756666	0.6269882	"protein phosphatase 1, catalytic subunit, alpha isoform"
810156	0.6266313	deoxythymidylate kinase (thymidylate kinase)
2015517	0.6259564	hypothetical protein FLJ22237
sd-71385		

Atty Dkt: 485772004300

770845	0.6253839	hexokinase 1
813707	0.6252347	regulator of G-protein signalling 16
611443	0.6244565	

01110	0.02 1 1000	myoglobii i
124331	0.624455	"cleavage and polyadenylation specific factor 5, 25 kD
127001	0.024400	cleavage and polyaderlylation specific factor 5, 25 kD
		subunit"
		Suburiil

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855800	0.6235647	prolyl endopeptidase
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625234	0.62188	KDEL (Lys-Asp-Glu-Leu) endoplasmic reticulum protein
		retention receptor 3

		i diamini i doopioi o
752272	0 6218222	hypothetical protein FLJ22649 similar to signal peptidase
133310	0.0210232	hypothetical protein FLJZZ049 similar to signal peptigase
		SPC22/23

		01 022/20
2055807	0.6216372	protein kinase domains containing protein similar to
		phosphoprotein C8FW

840878	0.6216124	seladin-1
1700436	0.6215741	FSTs

261472	0.6215341	putative nuclear	protein ORF1-FL49

2018821	0.6205344	ATPase inhibitor precursor
770700		DI/ETDE0404040

120749	0.6172281	"ESTs,	, Moderately similar to	KIAA1215	protein	[H.sapiens]
		,	industratory chilman to	> 1 (1) U (1 = 1)		[i i.oupiciio]

358162	0.616765	protein predicted by clone 23627
701212		les on a the attend of the total AAAA

781342	0.6159353	hypothetical protei	n MGC11115

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700000		s, Weakly similar to AAB47496 NG5 [H.sapiens]"
//nn44	0 6131441 "EST	S MASKIV similar to AARAMAG NCE IH canional"
1 20000	U.U IU I TU I	3, Weakly Sillillal to MAD4/430 NG3 II I.SableliSi
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2326057	0.61263	MLN51	protein
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701110	0.040004E	3.875 () · ·	
784140	ロ りつしノンダウ	VVI I reneat domain 1	~
701110	0.0102270	WD repeat domain 1	J

^{770355 0.6099761 &}quot;lanosterol synthase (2,3-oxidosqualene-lanosterol cyclase)"

^{81336 0.6099072} uteroglobin

^{757328 0.6097389} hypothetical protein FLJ22678

^{773674 0.6095732} oncogene TC21

^{0.6088919} ornithine decarboxylase antizyme 2

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826363 1758590		lysophospholipase II "fatty-acid-Coenzyme A ligase, long-chain 3"
1553306	0.6066787	"proteasome (prosome, macropain) 26S subunit, non- ATPase, 11"
785766	0.6058332	hypothetical protein
725841	0.6051304	KIAA0662 gene product
2020898	0.6044163	"procollagen-lysine, 2-oxoglutarate 5-dioxygenase 3"
504308	0.6041122	hypothetical protein FLJ10540
510575	0.6040713	hypothetical protein FLJ22087
49630	0.6027878	"calcium channel, voltage-dependent, L type, alpha 1D subunit"
142586	0.602037	MCT-1 protein
725284	0.6014099	"phosphorylase kinase, gamma 2 (testis)"
429799	0.6012713	hypothetical protein FLJ21939 similar to 5-azacytidine induced gene 2
1474424	0.6006702	"Homo sapiens cDNA FLJ12758 fis, clone NT2RP2001328"
784105	0.6003082	
2018527	0.599584	dolichyl-phosphate mannosyltransferase polypeptide 3
855563	0.5987092	v-erb-b2 avian erythroblastic leukemia viral oncogene homolog 3
823574	0.5978052	endosulfine alpha
323693	0.5975785	"adaptor-related protein complex 1, sigma 1 subunit"
824962	0.595868	"karyopherin alpha 2 (RAG cohort 1, importin alpha 1)"
773426	0.5941568	KIAA0391 gene product
772925		HSPCO34 protein
1656062	0.593408	coagulation factor XII (Hageman factor)
825740		DKFZp434J1813 protein
376516		cell division cycle 4-like
292213		"guanine nucleotide binding protein (G protein), beta polypeptide 2"
1898619	0.5880791	hypothetical protein MGC15737
414992		K562 cell-derived leucine-zipper-like protein 1
1573946	0.5875624	programmed cell death 9
739109		"adaptor-related protein complex 2, sigma 1 subunit"
1573946	0.5875624	programmed cell death 9

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430235 2572170		"H2B histone family, member Q" "ESTs, Weakly similar to T26581 hypothetical protein Y32B12A.3 - Caenorhabditis elegans [C.elegans]"
487733-2 1698036 127646 346696 74738	0.5843382 0.5838788 0.5838269	"NADH dehydrogenase (ubiquinone) 1 beta subcomplex, 2 (8kD, AGGG)" ubiquitin-conjugating enzyme E2 variant 1 TEA domain family member 4 "Homo sapiens, clone IMAGE:3535294, mRNA, partial cds"
124781 1883028		squalene epoxidase Homo sapiens mRNA; cDNA DKFZp434J1912 (from clone DKFZp434J1912)
183200	0.5801312	fumarylacetoacetate hydrolase (fumarylacetoacetase)
51741 590759 755301 843054 595213 752643 141852	0.5791873 0.5790795 0.5776102 0.5775511 0.5774681	GTP-binding protein sterol-C4-methyl oxidase-like "protein kinase C, delta" KIAA1533 protein hypothetical protein group XII secreted phospholipase A2 "purinergic receptor P2Y, G-protein coupled, 2"
813631 1601947		seven transmembrane protein TM7SF3 cytochrome c oxidase subunit VIIa polypeptide 2 (liver)
51657 814209 72050	0.5736866	hypothetical protein ET ESTs "chloride channel, nucleotide-sensitive, 1A"
1635665	0.5714089	"Homo sapiens, RIKEN cDNA 2010100012 gene, clone MGC:14813 IMAGE:4133274, mRNA, complete cds"
66406 530310 41356	0.5686743	hypothetical protein DKFZp762E1312 KIAA0143 protein "protein phosphatase 2, regulatory subunit B (B56), alpha isoform"
1844765	0.5670398	Homo sapiens mRNA; cDNA DKFZp564O1763 (from clone DKFZp564O1763)

489823	0.5666593	"COX17 (yeast) homolog, cytochrome c oxidase assembly protein"
1517749 813410	0.5665848 0.5664086	ESTs polymerase (RNA) II (DNA directed) polypeptide K (7.0kD)
1416782 1422338		"creatine kinase, brain" ribonucleotide reductase M2 polypeptide
250313 235986		"wingless-type MMTV integration site family, member 11"
66317 279720		"H1 histone family, member 2" "ESTs, Moderately similar to A47582 B-cell growth factor precursor [H.sapiens]"
884498	0.5640535	uncharacterized hypothalamus protein HT012
503851	0.564048	nuclear receptor co-repressor/HDAC3 complex subunit
823930	0.5633774	"actin related protein 2/3 complex, subunit 1A (41 kD)"
120271		hypothetical protein MGC4692 "interferon, alpha-inducible protein (clone IFI-6-16)"
782513		
246800		hypothetical protein FLJ10803
2309073		frizzled (Drosophila) homolog 5
784150		"RAB31, member RAS oncogene family"
139835	0.5587185	UDP-glucose dehydrogenase
1641894 796757		"adaptor-related protein complex 3, sigma 1 subunit"
813616		FK506-binding protein like
2549634		activator of S phase kinase
741769		"polymerase (DNA directed), beta"
488642	0.5560728	"ESTs, Weakly similar to I38022 hypothetical protein [H.sapiens]"
795498		putative transmembrane protein
1492780	0.5555908	"Homo sapiens cDNA FLJ14459 fis, clone HEMBB1002409"
241043		"Human clone 137308 mRNA, partial cds"
295986	0.5544422	emopamil-binding protein (sterol isomerase)
839682	0.5542571	ubiquitin-conjugating enzyme E2N (homologous to yeast UBC13)

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1581941 809466		hypothetical protein FLJ14540 DNA segment on chromosome 19 (unique) 1177 expressed sequence
1947381	0.5524634	hypothetical protein FLJ22329
825470	0.5523318	topoisomerase (DNA) II alpha (170kD)
1848977	0.5519913	glycerol kinase
700792	0.5493223	cyclin-dependent kinase inhibitor 3 (CDK2-associated dual specificity phosphatase)
825386	0.5487104	"ATP synthase, H+ transporting, mitochondrial F1F0, subunit d"
826194	0.5486839	synaptotagmin-like 2
299815	0.5485843	hypothetical protein DC42
1753497	0.5485231	ovo (Drosophila) homolog-like 1
744944	0.5482368	myosin VI
1499828	0.5479012	"fucosyltransferase 1 (galactoside 2-alpha-L- fucosyltransferase, Bombay phenotype included)"
415102	0.5465851	cell division cycle 25C
813387		diaphorase (NADH/NADPH) (cytochrome b-5 reductase)
786265		KIAA0750 gene product
1591264		transaldolase 1
756595	0.5433619	"S100 calcium-binding protein A10 (annexin II ligand, calpactin I, light polypeptide (p11))"
855749	0.5432466	triosephosphate isomerase 1
470092		like-glycosyltransferase
244764		B7 homolog 3
1637282		hexokinase 2
417801	0.5414761	mitochondrial ribosomal protein L27
826077	0.5405591	pyruvate dehydrogenase (lipoamide) beta
757489	0.5386267	"tubulin, alpha 2"
327635	0.5368033	adenylate kinase 1
343731	0.5365018	
786067	0.5361882	cell division cycle 25B
2017415	0.5353408	centromere protein A (17kD)
292996	0.5341429	"tyrosine 3-monooxygenase/tryptophan 5-monooxygenase activation protein, eta polypeptide"
815781	0.5340254	heat shock 105kD
346942		"phosphatidylinositol glycan, class Q"
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823694	0.5321203	"Homo sapiens chromosome 19, BAC CIT-HSPC_204F22 (BC228680), complete sequence; contains bacterial insertion element"
139354 221499 263727	0.5317861	hypothetical protein KIAA0508 protein "DNA segment, single copy probe LNS-CAI/LNS-CAII (deleted in polyposis"
489594	0.5308806	hypothetical protein FLJ11565
824524	0.5296138	UDP-galactose transporter related
951216	0.5291832	"NADH dehydrogenase (ubiquinone) 1 beta subcomplex, 10 (22kD, PDSW)"
841260	0.5285587	hypothetical protein
283739	0.5276555	"Homo sapiens cDNA FLJ14028 fis, clone HEMBA1003838"
49117		KIAA0215 gene product
1762111	0.5272967	natriuretic peptide receptor C/guanylate cyclase C (atrionatriuretic peptide receptor C)
358609	0.5266332	NADH dehydrogenase (ubiquinone) flavoprotein 3 (10kD)
884425	0.5265114	"chaperonin containing TCP1, subunit 5 (epsilon)"
795401	0.5264278	diacylglycerol O-acyltransferase (mouse) homolog
811907	0.5246416	hypothetical protein FLJ22056
1631747		male-enhanced antigen
365060		"RAB11A, member RAS oncogene family"
743220		hypothetical protein FLJ12517
705064	0.5238148	"transforming, acidic coiled-coil containing protein 3"
768452	0.5237494	"Homo sapiens EST from clone 491476, full insert"
743977	0.5237158	Homo sapiens mRNA for TL132
1568825	0.5205165	Arg/Abl-interacting protein ArgBP2
772898	0.520392	ribosomal protein S15a
366834	0.5191005	envoplakin
1616253		breast carcinoma amplified sequence 1
322617	0.5188645	v-ral simian leukemia viral oncogene homolog B (ras related; GTP binding protein)
825296	0.5178447	low density lipoprotein receptor defect C complementing

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491524 143426 73009	0.5173212	mitochondrial ribosomal protein L13 "ras homolog gene family, member B" "ESTs, Weakly similar to A43932 mucin 2 precursor, intestinal [H.sapiens]"
293569	0.5162069	chromosome 1 open reading frame 21
296702		"deiodinase, iodothyronine, type I"
773286	0.5158979	"solute carrier family 9 (sodium/hydrogen exchanger),
		isoform 3 regulatory factor 1"
726658	0.5157542	"non-metastatic cells 3, protein expressed in"
753897	0.5153412	autocrine motility factor receptor
293727		hypothetical protein MGC861
810947	0.5151346	"LIS1-interacting protein NUDE1, rat homolog"
491465	0.5150527	hypothetical protein FLJ10035
814899		BCL2/adenovirus E1B 19kD-interacting protein 3-like
000000	0 5405550	ECT-
266500 1159963	0.5135558	interferon regulatory factor 7
785707		protein regulator of cytokinesis 1
346134		calcium-regulated heat-stable protein (24kD)
486110	0.5127183	•
2017403		regulator of G-protein signalling 3
491527	0.5116183	"Homo sapiens, Similar to CG7083 gene product, clone MGC:10534 IMAGE:3957147, mRNA, complete cds"
		WIGO. 10004 IMAGE. 3907 147, MINNA, Complete cus
769600		uracil-DNA glycosylase 2
1558233	0.5101529	
649084		carbonic anhydrase XI
825822 1461477		hypothetical protein
1401477	0.509551	Homo sapiens mRNA; cDNA DKFZp586I0324 (from clone DKFZp586I0324)
		•
327506	0.5092192	Homo sapiens mRNA full length insert cDNA clone
769942	0.509086	EUROIMAGE 327506 kinesin-like 4
505289		"angiotensin II, type I receptor-associated protein"
726439		CGI-143 protein
79726	0.50/86//	"ESTs, Highly similar to T46395 hypothetical protein
		DKFZp434I1120.1 [H.sapiens]"

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810762 785793		SNARE protein "capping protein (actin filament) muscle Z-line, alpha 1"
81599	0.5067033	ubiquitin specific protease 14 (tRNA-guanine transglycosylase)
294397	0.5066683	DKFZP586A0522 protein
1947647		CGI-147 protein
795805		KIAA0332 protein
810959	0.505619	Rho GDP dissociation inhibitor (GDI) alpha
2322223	0.5047814	small nuclear ribonucleoprotein polypeptide A
472103	0.503987	"soc-2 (suppressor of clear, C.elegans) homolog"
810609		hypothetical protein PP1226
897813	0.5034689	polyadenylate binding protein-interacting protein 1
1500542 839746	0.5032679 0.5031787	regulator of G-protein signalling 11 "Homo sapiens, Similar to RIKEN cDNA 5830420C20 gene, clone IMAGE:3633379, mRNA, partial cds"
343607 649977	0.5029133 0.5002695	AD-015 protein Homo sapiens clone CDABP0014 mRNA sequence
823940	0 4997308	"transducer of ERBB2, 1"
2250839		androgen receptor (dihydrotestosterone receptor; testicular feminization; spinal and bulbar muscular atrophy; Kennedy disease)
731080	0.4979864	hypothetical protein FLJ12661
753400	0.496254	CGI-204 protein
869375	0.4958886	"isocitrate dehydrogenase 2 (NADP+), mitochondrial"
454040	0.4040574	dynastin 1
154610		dynactin 4 "Homo sapiens, Similar to clone FLB3816, clone
130835	0.4940763	IMAGE:3454380, mRNA"
859228	0.4942316	"isocitrate dehydrogenase 1 (NADP+), soluble"
41698		progesterone binding protein
30170	0.4936252	"caspase 3, apoptosis-related cysteine protease"
266218	0.4934791	hypothetical protein FLJ11350

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509588	0.49319	"TATA box binding protein (TBP)-associated factor, RNA polymerase II, J, 20kD"
1916575	0.4923447	BCL2-interacting killer (apoptosis-inducing)
744994 487444 503866 814353	0.4887966 0.4884193	hypothetical protein FLJ12242 "cyclic AMP phosphoprotein, 19 kD" sperm autoantigenic protein 17 phorbol-12-myristate-13-acetate-induced protein 1
815501 746163		hypothetical protein MGC2721 "ESTs, Weakly similar to ALU1_HUMAN ALU SUBFAMILY J SEQUENCE CONTAMINATION WARNING ENTRY [H.sapiens]"
347726	0.4837701	homeo box D8
897770	0.4836347	
1854539	0.4824023	GAP-associated tyrosine phosphoprotein p62 (Sam68)
41123	0.481642	"Homo sapiens, Similar to RIKEN cDNA 2210021G21 gene, clone MGC:14859 IMAGE:3621871, mRNA, complete cds"
856447	0.4816395	"interferon, gamma-inducible protein 30"
1568561		BCL2-like 1
744047		polo (Drosophia)-like kinase
290101	0.4800598	
455275		hypothetical protein FLJ23469
2069602	0.4786416	melanocortin 1 receptor (alpha melanocyte stimulating hormone receptor)
592801	0.4785497	"serine palmitoyltransferase, long chain base subunit 2"
741891	0.4775456	"RAB2, member RAS oncogene family-like"
1434948	0.4771454	HIV TAT specific factor 1
430614		"2,3-bisphosphoglycerate mutase"
454896		"DnaJ (Hsp40) homolog, subfamily A, member 2"
268946	0.4751692	WD40 protein Ciao1
1572710		hypothetical protein FLJ21213
626318		ubinuclein 1
377384	0.4746085	"nuclear receptor subfamily 2, group F, member 2"

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1632252	0.474559	"complement component 1, q subcomponent, alpha polypeptide"
489444	0.4741367	hypothetical protein FLJ20211
345787		"highly expressed in cancer, rich in leucine heptad repeats"
767487	0.4729158	"ariadne (Drosophila) homolog, ubiquitin-conjugating enzyme E2-binding protein, 1"
empty-31	0.4726918	blank
135640	0.4708685	syntaxin 3A
754653	0.4707927	"cadherin, EGF LAG seven-pass G-type receptor 3, flamingo (Drosophila) homolog"
1557637	0.4705061	ESTs
788641	0.4703576	"adaptor-related protein complex 1, sigma 2 subunit"
810567	0.4702908	"Homo sapiens, clone MGC:3182 IMAGE:3356293, mRNA, complete cds"
366039	0.4701924	KIAA0892 protein
781047	0.4699376	budding uninhibited by benzimidazoles 1 (yeast homolog)
782688	0.4694868	"dynein, axonemal, light intermediate polypeptide"
210862	0.4692392	"acyl-Coenzyme A oxidase 1, palmitoyl"
510794	0.4691467	c-myc binding protein
121436	0.4691298	"Homo sapiens, clone MGC:4677 IMAGE:3532809, mRNA, complete cds"
325515	0.4689353	hypothetical protein FLJ10980
965223		"thymidine kinase 1, soluble"
1469148	0.467634	FGFR1 oncogene partner
149539	0.4671676	KIAA1700
2306987	0.4670155	secreted and transmembrane 1
122147	0.4652151	
2063982	0.4651856	"potassium channel, subfamily K, member 6 (TWIK-2)"
769890	0.4647115	nucleoside phosphorylase
1642496	0.4645092	hypothetical protein MGC11266
428582	0.4642013	hypothetical protein FLJ20296
66599	0.4639123	N-acetyltransferase 1 (arylamine N-acetyltransferase)
1565455	0.4634802	"nuclear receptor subfamily 2, group C, member 2"
810939	0.4634658	hypothetical protein FLJ22169
359723		mitochondrial elongation factor G
ad 71295		

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1027283 1636844 469151		ESTs ring finger protein 14 "eukaryotic translation initiation factor 2, subunit 2 (beta, 38kD)"
813675	0.4614894	"Human D9 splice variant B mRNA, complete cds"
489657 469383 40042 726637	0.4607375 0.4604322	tryptophan rich basic protein chromosome 8 open reading frame 1 hypothetical protein FLJ10747 t-complex-associated-testis-expressed 1-like
811142 179212		"phosphoinositide-3-kinase, regulatory subunit, polypeptide 2 (p85 beta)" "ESTs, Moderately similar to T12539 hypothetical protein
292936 2046679	0.4593712 0.4579596	DKFZp434J154.1 [H.sapiens]" hypothetical protein FLJ10468
810497	0.4572069	"ESTs, Weakly similar to A35363 synapsin I splice form a [H.sapiens]"
1585492 782689	0.4571478 0.4570254	"solute carrier family 6 (neurotransmitter transporter, creatine), member 8"
774446 366156	• • • • • • • • • • • • • • • • • • • •	adrenomedullin "Homo sapiens cDNA FLJ14028 fis, clone HEMBA1003838"
725649	0.4559473	"nuclear factor of activated T-cells, cytoplasmic, calcineurin-dependent 4"
76362	0.4553121	"spectrin, alpha, non-erythrocytic 1 (alpha-fodrin)"
754046	0.4550413	DNA segment on chromosome X (unique) 9879 expressed sequence
345538 782439		cathepsin L "ATP synthase, H+ transporting, mitochondrial F0 complex, subunit e"
1537001 345423 1702742		
825659 1652310		system), member 5" N-myc downstream regulated "Homo sapiens, clone MGC:19613 IMAGE:3833049,
ad 71205		mRNA, complete cds"

40773 0.4510289 "guanine nucleotide binding protein (G protein), alpha z polypeptide" 35626 0.4509825 "Homo sapiens cDNA FLJ14201 fis, clone NT2RP3002955" 358217 0.4507923 glypican 4 214996 0.45033 ESTs 177827 0.4500254 synaptotagmin VII 1075635 0.4495618 MLSN1- and TRP-related 810039 0.4495188 defender against cell death 1 772220 0.4475721 for protein disulfide isomerase-related 70606 0.4475721 smg GDS-ASSOCIATED PROTEIN 841621 0.4466003 "ESTs, Weakly similar to JE0350 Anterior gradient-2 [H.sapiens]" 855707 0.4456433 ESTs 1455394 0.4456695 cytochrome c 85804 0.44449 hypothetical protein FLJ21918 809512 0.4442565 hypothetical protein FLJ10767 753457 0.4435792 NADH dehydrogenase (ubiquinone) Fe-S protein 1 (75kD) (NADH-coenzyme Q reductase) 767289 0.4432685 hypothetical protein FLJ10055 565235 0.4428854 spermine synthase 611467 0.4424976 ALL1-fused gene from chromosome 1q 1558642 0.4424721 hypothetical protein MGC2771 814989 0.4419384 "protein phosphatase 1G (formerly 2C), magnesium-dependent, gamma isoform" 1899312 0.4419344 keratin 16 (focal non-epidermolytic palmoplantar keratoderma) 1565079 0.441936 B-cell linker 783681 0.4411931 interferon-induced protein 35 785368 0.441074 PDZ-binding kinase; T-cell originated protein kinase 1517595 0.4400759 IKIAAO175 gene product 713862 0.4400704 ubiquitin-protein isopeptide ligase (E3)	1523225	0 4511104	oncostatin M receptor
358217 0.4509825 "Homo sapiens cDNA FLJ14201 fis, clone NT2RP3002955" 358217 0.4507923 glypican 4 214996 0.45033 ESTs 177827 0.4500254 synaptotagmin VII 1075635 0.4495818 MLSN1- and TRP-related 810039 0.4491986 defender against cell death 1 772220 0.4475721 for protein disulfide isomerase-related 76060 0.4475892 ESTs 1692195 0.4471181 smg GDS-ASSOCIATED PROTEIN 841621 0.4466003 "ESTs, Weakly similar to JE0350 Anterior gradient-2 [H.sapiens]" 855707 0.4456433 ESTs 1455394 0.445565 cytochrome c 85804 0.44449 hypothetical protein FLJ21918 809512 0.4442565 hypothetical protein FLJ10767 753457 0.4435792 NADH dehydrogenase (ubiquinone) Fe-S protein 1 (75kD) (NADH-coenzyme Q reductase) 767289 0.4432685 hypothetical protein FLJ10055 565235 0.4428854 spermine synthase 611467 0.4428543 NADH dehydrogenase (ubiquinone) Fe-S protein 6 (13kD) (NADH-coenzyme Q reductase) 812105 0.4428854 spermine synthase 611467 0.4424761 ALL1-fused gene from chromosome 1q 15558642 0.442721 hypothetical protein MGC2771 814989 0.441934 "protein phosphatase 1G (formerly 2C), magnesium-dependent, gamma isoform" 1899312 0.4418248 ESTs 1486533 0.441804 keratin 16 (focal non-epidermolytic palmoplantar keratoderma) 1565079 0.4415966 B-cell linker 783681 0.4411742 PDZ-binding kinase; T-cell originated protein kinase 1517595 0.4409591 KIAA0175 gene product 713862 0.4400704 ubiquitin-protein isopeptide ligase (E3)			"guanine nucleotide binding protein (G protein), alpha z
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214996 0.45033 ESTs 177827 0.4500254 synaptotagmin VII 1075635 0.4491986 defender against cell death 1 772220 0.4475721 for protein disulfide isomerase-related 70606 0.4475592 ESTs 1692195 0.4471181 smg GDS-ASSOCIATED PROTEIN 841621 0.4466003 "ESTs, Weakly similar to JE0350 Anterior gradient-2 [H.sapiens]" 855707 0.4456935 ESTs 1455394 0.4455935 cytochrome c 85804 0.44449 hypothetical protein FLJ21918 809512 0.4442565 hypothetical protein FLJ10767 753457 0.4432685 hypothetical protein FLJ100767 767289 0.4432685 hypothetical protein FLJ10055 565235 0.4428543 NADH dehydrogenase (ubiquinone) Fe-S protein 6 (13kD) (NADH-coenzyme Q reductase) 812105 0.4428543 NADH dehydrogenase (ubiquinone) Fe-S protein 6 (13kD) (NADH-coenzyme Q reductase) 812105 0.4424976 ALL1-fused gene from chromosome 1q 1558642 0.4419384 "protein phosphatase 1G (formerly 2C), magnesium-depen	33323	0000020	•
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keratoderma) 1565079 0.4415966 B-cell linker 783681 0.4412503 upstream regulatory element binding protein 1 1422723 0.4411931 interferon-induced protein 35 785368 0.4411742 PDZ-binding kinase; T-cell originated protein kinase 1517595 0.4409591 KIAA0175 gene product 713862 0.4400704 ubiquitin-protein isopeptide ligase (E3)	1899312	0.4418248	ESTs
1565079 0.4415966 B-cell linker 783681 0.4412503 upstream regulatory element binding protein 1 1422723 0.4411931 interferon-induced protein 35 785368 0.4411742 PDZ-binding kinase; T-cell originated protein kinase 1517595 0.4409591 KIAA0175 gene product 713862 0.4400704 ubiquitin-protein isopeptide ligase (E3)	1486533	0.441804	
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1517595 0.4409591 KIAA0175 gene product 713862 0.4400704 ubiquitin-protein isopeptide ligase (E3)	1422723	0.4411931	interferon-induced protein 35
713862 0.4400704 ubiquitin-protein isopeptide ligase (E3)	785368	0.4411742	PDZ-binding kinase; T-cell originated protein kinase
713862 0.4400704 ubiquitin-protein isopeptide ligase (E3)	1517595	0.4409591	KIAA0175 gene product
	sd-71385		

83358 81203 74070 2306221	0.4390933	ESTs paraoxonase 3 endosulfine alpha "wingless-type MMTV integration site family, member 10B"
150118 214068 1434905 450301	0.4383007 0.4382073 0.4380657 0.437854	ESTs hypothetical protein DKFZp434F054 GATA-binding protein 3 homeo box B7 mutL (E. coli) homolog 3 hypothetical protein FLJ20297 "Homo sapiens cDNA: FLJ23249 fis, clone COL04196"
824108 795543		SCAN domain-containing 1 thioredoxin peroxidase (antioxidant enzyme)
1592837	0.4355927	interferon induced transmembrane protein 2 (1-8D)
77533	0.435582	"inositol polyphosphate-5-phosphatase, 40kD"
1583198	0.4350835	"ESTs, Weakly similar to S65824 reverse transcriptase homolog [H.sapiens]"
84295	0.4349213	interleukin 1 receptor antagonist
770992	0.4346858	
837864	0.4345283	progestin induced protein
1909574	0.4342055	mitochondrial ribosomal protein S11
128695	0.4340119	"ESTs, Weakly similar to I38344 titin, cardiac muscle [H.sapiens]"
742581	0.433453	"Homo sapiens cDNA FLJ10366 fis, clone NT2RM2001420"
52103	0.432756	hypothetical protein FLJ23045
770675	0.4319407	"Homo sapiens cDNA: FLJ21323 fis, clone COL02374"
756373	0.4310628	Rho guanine exchange factor (GEF) 16
43977	0.4309655	KIAA0182 protein
2322079	0.4309341	
25440	0.4304195	"staufen (Drosophila, RNA-binding protein) homolog 2"
207794	0.4297389	"nuclear factor (erythroid-derived 2), 45kD"
746190	0.4291743	hypothetical protein DKFZp761B1514
74566		"exportin 1 (CRM1, yeast, homolog)"
76196	0.4278385	hypothetical protein FLJ20062
nd 71205		

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826173 1587863	0.4276784 0.4270229	profilin 1 acetyl-Coenzyme A acyltransferase 1 (peroxisomal 3- oxoacyl-Coenzyme A thiolase)
2017721	0.4268089	"ESTs, Weakly similar to 2109260A B cell growth factor [H.sapiens]"
809464	0.426456	"fibroblast growth factor receptor 2 (bacteria-expressed kinase, keratinocyte growth factor receptor, craniofacial dysostosis 1, Crouzon syndrome, Pfeiffer syndrome, Jackson-Weiss syndrome)"
41826 1687976	0.4264531 0.4257058	ESTs "endoplasmic reticulum chaperone SIL1, homolog of yeast"
416436 145491 810316	0.4250102	mitochondrial ribosomal protein L24 protocadherin 1 (cadherin-like 1) very long-chain acyl-CoA synthetase; lipidosin
629944 1637302 754625 298417 47853	0.424164 0.4236648 0.4235025	myosin VB DNAJ domain-containing "ATPase, Class II, type 9A" trefoil factor 3 (intestinal) "aldehyde dehydrogenase 4 family, member A1"
809727 774502		unc-51 (C. elegans)-like kinase 1 "protein tyrosine phosphatase, non-receptor type 12"
785840	0.4219115	"SEC24 (S. cerevisiae) related gene family, member D"
823907 1679942 823909 1600239	0.4215618 0.4215098 0.421459	hypothetical protein FLJ10511 KIAA1053 protein HSPC037 protein
85195	0.4207753	"growth arrest and DNA-damage-inducible, gamma"
1640282 782503 266312	0.4205398 0.4201146	KIAA1240 protein fatty acid desaturase 1 "ATPase, Cu++ transporting, beta polypeptide (Wilson disease)"
884511 2051697		cytochrome c oxidase subunit VIIb "inhibitor of kappa light polypeptide gene enhancer in B- cells, kinase beta"

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724888	0.4196322	"cytochrome P450, subfamily IVB, polypeptide 1"
809507 151449		hypothetical protein FLJ20568 "protein tyrosine phosphatase, non-receptor type 21"
743860 345833		"hypothetical protein, clone 2746033" heterogeneous nuclear ribonucleoprotein A/B
259950 283751 810875 26883	0.4179154 0.4176896	hypothetical protein FLJ14991 cortistatin "solute carrier family 26, member 6" "protein kinase (cAMP-dependent, catalytic) inhibitor beta"
38356 35191		"cytochrome P450, subfamily 46 (cholesterol 24- hydroxylase)" stromal cell-derived factor 2
75859 1569187		N-myc downstream-regulated gene 2 heparan sulfate (glucosamine) 3-O-sulfotransferase 4
345670	-1.889471	"ESTs, Moderately similar to I59348 CCAAT binding transcription factor CBF subunit C - rat [R.norvegicus]"
760299 753071		dickkopf (Xenopus laevis) homolog 3 "Homo sapiens cDNA: FLJ22528 fis, clone HRC12825"
344720 810002		glycophorin C (Gerbich blood group) "Homo sapiens, clone MGC:19762 IMAGE:3636045, mRNA, complete cds"
200814	-1.7140538	"membrane metallo-endopeptidase (neutral endopeptidase, enkephalinase, CALLA, CD10)"
131839 300632		folate receptor 1 (adult) hypothetical protein FLJ21044 similar to Rbig1
811920 486683		"interleukin 11 receptor, alpha" Homo sapiens mRNA; cDNA DKFZp564J0323 (from clone DKFZp564J0323)
1558675 840266		SRY (sex determining region Y)-box 10 "Homo sapiens cDNA: FLJ22667 fis, clone HSI08385"
285377 1469377		pellino (Drosophila) homolog 2 lipoma HMGIC fusion partner-like 2

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45099	-1.6325981	regucalcin (senescence marker protein-30)
1161564 298122 796542 289760	-1.5553964	desmuslin frizzled (Drosophila) homolog 7 ets variant gene 5 (ets-related molecule) "ESTs, Highly similar to T00391 hypothetical protein KIAA0612 [H.sapiens]"
22917	-1.5156635	Homo sapiens mRNA; cDNA DKFZp761M0111 (from clone DKFZp761M0111)
838478	-1 5059801	neurocalcin delta
788234		"inhibitor of DNA binding 4, dominant negative helix-loophelix protein"
377461	-1.4787671	"caveolin 1, caveolae protein, 22kD"
67741	-1.4786722	PP2135 protein
839736		"crystallin, alpha B"
712139		ADP-ribosylation factor-like 7
160192		"ESTs, Weakly similar to 2004399A chromosomal protein [H.sapiens]"
742685	-1.4440371	disabled (Drosophila) homolog 2 (mitogen-responsive phosphoprotein)
1556433	-1.4191955	GRO3 oncogene
1554167	-1.4126653	hypothetical protein FLJ14529
877621		nGAP-like protein
1882697		peanut (Drosophila)-like 2
72778	-1.3873177	"caspase 7, apoptosis-related cysteine protease"
666879	-1.3624072	annexin A8
69002	-1.3581187	PPAR(gamma) angiopoietin related protein
811848	-1.3549757	hypothetical protein
490023	-1.3516302	hypothetical protein MGC2648
781017	-1.3232108	early growth response 2 (Krox-20 (Drosophila) homolog)
767202	-1.3217103	latent transforming growth factor beta binding protein 2
291478		runt-related transcription factor 3
757191	-1.2902603	intercellular adhesion molecule 2
130201 1635320		"amiloride-sensitive cation channel 2, neuronal"

140574	-1.2754214	"small inducible cytokine subfamily D (Cys-X3-Cys), member 1 (fractalkine, neurotactin)"
416676 814798		pellino (Drosophila) homolog 1 "aldehyde dehydrogenase 1 family, member A3"
611532 2056139 377275	-1.2697411	"troponin I, skeletal, fast" LIM domain protein ataxia-telangiectasia group D-associated protein
303109 813265		purinergic receptor (family A group 5) Homo sapiens mRNA; cDNA DKFZp564H1916 (from clone DKFZp564H1916)
878836	-1.2507242	"secretory granule, neuroendocrine protein 1 (7B2 protein)"
1473471 529843		KIAA0194 protein "ESTs, Moderately similar to JC5238 galactosylceramide-like protein, GCP [H.sapiens]"
505864 781014 470393	-1.2359239	RalGDS-like gene suppression of tumorigenicity 5 "matrix metalloproteinase 7 (matrilysin, uterine)"
290378 196435 71087	-1.2282733 -1.2181178	"v-maf musculoaponeurotic fibrosarcoma (avian) oncogene family, protein F"
76182	-1.2102812	hypothetical protein DKFZp761F241
884462		Down syndrome critical region gene 1 KIAA1706 protein
277571 841308		"myosin, light polypeptide kinase"
162308	-1.2006293	
52419		Friedreich ataxia region gene X123
488404	-1.1981268	Homo sapiens clone TUA8 Cri-du-chat region mRNA
712401	-1.193113	"phosphoinositide-3-kinase, catalytic, delta polypeptide"
289428	-1.1923798	"neurotrophic tyrosine kinase, receptor, type 2"
814443 153760	-1.1905978 -1.1793765	hypothetical protein MGC3232 EphB1
1 71205		

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1917449 1493218 843070 753038 811088 2106144 796181 1257131 1473274	-1.164886 -1.1647509 -1.1629978 -1.1625002 -1.1619925 -1.1584587 -1.1566479	regulated in glioma growth arrest-specific 6
208718 811837	-1.1498532 -1.1468554	annexin A1 eukaryotic translation elongation factor 1 alpha 1
768571 1474900 756708	-1.1347123	SRY (sex determining region Y)-box 8 keratin 15 "potassium intermediate/small conductance calcium-activated channel, subfamily N, member 4"
1555924 811837 1878409 786069 815737	-1.1269309 -1.1144369 -1.1143906	CSR1 protein catechol-O-methyltransferase beta-site APP-cleaving enzyme "ATP synthase, H+ transporting, mitochondrial F1 complex, alpha subunit, isoform 1, cardiac muscle"
132857	-1.0989746	Homo sapiens mRNA; cDNA DKFZp586N1323 (from clone DKFZp586N1323)
1871116	-1.0975491	Homo sapiens mRNA; cDNA DKFZp434C1714 (from clone DKFZp434C1714); partial cds
133236	-1.0945253	"RNA binding motif protein, X chromosome"
283023 2016775		chemokine (C-X3-C) receptor 1 "G protein-coupled receptor, family C, group 5, member B"
897963 814316 47043 625399	-1.0654718 -1.0559861	phosphatidic acid phosphatase type 2A ribosomal protein L13 tensin hypothetical protein similar to beta-transducin family
416959 366100		nuclear factor I/B matrilin 2
sd-71385		

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753162	-1.0416438	KIAA0603 gene product
491403	-1.0411657	"tumor necrosis factor receptor superfamily, member 1B"
813256	-1.0407652	"ATP-binding cassette, sub-family B (MDR/TAP), member 1"
740620	-1.0395569	tropomyosin 2 (beta)
1577736	-1.035115	epidermal growth factor (beta-urogastrone)
70245	-1.0348655	Homo sapiens mRNA full length insert cDNA clone EUROIMAGE 50374
1626996	-1.0328113	c-fos induced growth factor (vascular endothelial growth factor D)
360254	-1.0325653	"cysteine-rich, angiogenic inducer, 61"
80344	-1.0323882	interleukin 7 receptor
530958	-1.0316423	smoothened (Drosophila) homolog
34150	-1.0302576	ESTs
809784	-1.030253	"kallikrein 6 (neurosin, zyme)"
990881	-1.0289558	
416434	-1.0284229	SCN Circadian Oscillatory Protein (SCOP)
162308	-1.023071	
796475	-1.0217348	four and a half LIM domains 3
32489	-1.0196444	hypothetical protein DKFZp566A1524
2504881	-1.0194516	signal transducer and activator of transcription 5A
188335	-1.0161259	"egf-like module containing, mucin-like, hormone receptor-like sequence 2"
593023	-1.0135883	"dystrobrevin, beta"
139660	-1.0134677	
772913	-1.0122192	calreticulin
270826	-1.0112851	"Homo sapiens cDNA FLJ13329 fis, clone OVARC1001795"
1864302	-1.0099902	E74-like factor 5 (ets domain transcription factor)
		period (Drosophila) homolog 1 Homo sapiens mRNA; cDNA DKFZp586M2022 (from clone DKFZp586M2022)
788136	-1.0073983	"phosphodiesterase 4B, cAMP-specific (dunce (Drosophila)-homolog phosphodiesterase E4)"
40027	-1.0057005	ESTs
1856063	-1.0036618	tweety (Drosophila) homolog 1
		zinc finger protein 221
sd-71385		

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	628955 2095066		forkhead box O1A (rhabdomyosarcoma) "alcohol dehydrogenase 1C (class I), gamma polypeptide"
	1636166 882248	-0.9969091 -0.9965858	KIAA0668 protein
	415816	-0.9954308	•
	897731	-0.9927981	
:	868396		"tyrosine 3-monooxygenase/tryptophan 5-monooxygenase activation protein, eta polypeptide"
	343760	-0.9907562	SH3 domain binding glutamic acid-rich protein like 2
	1572298	-0.9906095	"CD3Z antigen, zeta polypeptide (TiT3 complex)"
	1474337	-0.986117	"phosphorylase, glycogen; brain"
	1609665		BarH-like homeobox 2
	2243051	-0.9835274	kallikrein 8 (neuropsin/ovasin)
	27544	-0.9834143	prominin (mouse)-like 1
	281190	-0.9830533	ESTs
	161456	-0.9820939	serum amyloid A1
	505243	-0.980381	"inositol 1,4,5-triphosphate receptor, type 2"

Example IV: Genes for discriminating between normal and ADH

As shown in Table 3 below, 600 genes were identified as being able to discriminate between normal and ADH.

Table 3

ClonelD	Weight	Description
1323448	1.3237504	cysteine-rich protein 1 (intestinal)
594500	1.158305	EST
824052	1.034408	chromosome 6 open reading frame 1
149539	0.9878366	KIAA1700
788654	0.9585887	growth factor receptor-bound protein 2
1517749	0.947675	ESTs
1492238	0.9155471	HSPC003 protein
cd_71385		

sd-71385

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366353 1609836		DKFZP564C186 protein glutamate-ammonia ligase (glutamine synthase)
138189 344959 1435862	0.8945717	Wolfram syndrome 1 (wolframin) gene for serine/threonine protein kinase antigen identified by monoclonal antibodies 12E7, F21 and O13
2014373 23776 293569 130835	0.879215 0.8708985 0.8659688	HNK-1 sulfotransferase quinoid dihydropteridine reductase chromosome 1 open reading frame 21 Homo sapiens, Similar to clone FLB3816, clone IMAGE:3454380, mRNA uncoupling protein 2 (mitochondrial, proton carrier)
488025 178805	0.8525751	Kruppel-associated box protein Human DNA sequence from clone RP5-850E9 on chromosome 20. Contains part of the gene for a novel C2H2 type zinc finger protein similar to Drosophila Scratch (Scrt), Slug and Xenopus Snail, a novel gene similar to Drosophila CG6762, STSs, GSSs and five CpG
745606 1492463 530197	0.8378401	hypothetical protein PP591 selenoprotein X, 1 ESTs, Moderately similar to ALU8_HUMAN ALU SUBFAMILY SX SEQUENCE CONTAMINATION WARNING ENTRY [H.sapiens]
1492463	0.8378401 0.8206765	selenoprotein X, 1 ESTs, Moderately similar to ALU8_HUMAN ALU SUBFAMILY SX SEQUENCE CONTAMINATION WARNING ENTRY
1492463 530197 898222 782608 450301	0.8378401 0.8206765 0.8092364 0.8041049 0.8030667 0.7992668 0.7924672 0.7904354	selenoprotein X, 1 ESTs, Moderately similar to ALU8_HUMAN ALU SUBFAMILY SX SEQUENCE CONTAMINATION WARNING ENTRY [H.sapiens]
1492463 530197 898222 782608 450301 196189 363144 68636	0.8378401 0.8206765 0.8092364 0.8041049 0.8030667 0.7992668 0.7924672 0.7904354 0.7765346	selenoprotein X, 1 ESTs, Moderately similar to ALU8_HUMAN ALU SUBFAMILY SX SEQUENCE CONTAMINATION WARNING ENTRY [H.sapiens] Homo sapiens clone 24418 mRNA sequence mitochondrial ribosomal protein L9 mutL (E. coli) homolog 3 cytochrome b-5 transcription factor AP-2 beta (activating enhancer-binding protein 2 beta) hypothetical protein MGC2477

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1558233 769600 1156538	0.763743	ESTs uracil-DNA glycosylase 2 potassium inwardly-rectifying channel, subfamily J, member 11
1587863		acetyl-Coenzyme A acyltransferase 1 (peroxisomal 3-oxoacyl-
		Coenzyme A thiolase)
1709791 741891 788654		BAI1-associated protein 1 RAB2, member RAS oncogene family-like
-	0.7223819	CGI-49 protein a disintegrin-like and metalloprotease (reprolysin type) with thrombospondin type 1 motif, 4
120138 2248488		J domain containing protein 1 ems1 sequence (mammary tumor and squamous cell carcinoma-associated (p80/85 src substrate)
265103	0.702883	Homo sapiens mRNA; cDNA DKFZp547M123 (from clone DKFZp547M123)
2054122	0.698129	solute carrier family 11 (proton-coupled divalent metal ion transporters), member 3
701751	0.6961443	cut (Drosophila)-like 1 (CCAAT displacement protein)
811582 266312		golgi phosphoprotein 2 ATPase, Cu++ transporting, beta polypeptide (Wilson disease)
35147	0.6842172	ESTs, Weakly similar to unnamed protein product [H.sapiens]
1325816	0.6829962	polymerase (RNA) II (DNA directed) polypeptide L (7.6kD)
358217	0.6801945	
233349		hypothetical protein FLJ10761
741977		B-factor, properdin
145132	-	mannose-P-dolichol utilization defect 1
33267	0.6679827	TOT MAIL LANGUAGE A 47500 D cell amountly footon
279720	0.6607182	ESTs, Moderately similar to A47582 B-cell growth factor precursor [H.sapiens]
565319	0.6603668	Homo sapiens mRNA; cDNA DKFZp564B1264 (from clone DKFZp564B1264)
771173	0.6594128	mitochondrial ribosomal protein S21

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0.6581316 0.6563238	C3HC4-like zinc finger protein hypothetical protein FLJ11280 carnitine acetyltransferase transmembrane 7 superfamily member 1 (upregulated in kidney)
0.654954	androgen receptor (dihydrotestosterone receptor; testicular feminization; spinal and bulbar muscular atrophy; Kennedy disease)
0.6499845	ESTs, Moderately similar to T12539 hypothetical protein DKFZp434J154.1 [H.sapiens]
	RAB31, member RAS oncogene family Homo sapiens, clone IMAGE:3604869, mRNA
0.6440792 0.6399293	pilin-like transcription factor hypothetical protein FLJ22353 nuclear receptor coactivator 3 uncharacterized hematopoietic stem/progenitor cells protein MDS030
0.6388187	polymerase (DNA directed), beta hypothetical protein FLJ23403 Homo sapiens cDNA: FLJ22139 fis, clone HEP20959
0.630827	cyclin-dependent kinase 5 growth factor, erv1 (S. cerevisiae)-like (augmenter of liver regeneration) Homo sapiens mRNA; cDNA DKFZp434E2321 (from clone
	DKFZp434E2321); partial cds
0.6283163	calcium and integrin binding protein (DNA-dependent protein kinase interacting protein)
0.6268883	Homo sapiens cDNA: FLJ21587 fis, clone COL06946
	mitochondrial ribosomal protein L23 1-acylglycerol-3-phosphate O-acyltransferase 2 (lysophosphatidic acid acyltransferase, beta)
0.6205287 0.6203153	citrate synthase synaptotagmin-like 2 artemin hypothetical protein FLJ22056
	0.6581316 0.6563238 0.6554059 0.6554059 0.64954 0.6499845 0.6495706 0.6457259 0.6440792 0.6399293 0.6398576 0.6390692 0.6388187 0.6329458 0.6329458 0.6299822 0.6299822 0.6283163

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814350 611443 782608	0.6173648 0.6112713 0.6105794	insulin-degrading enzyme myoglobin
22778 128695		sulfortranferase family 4A, member 1 ESTs, Weakly similar to l38344 titin, cardiac muscle [H.sapiens]
47853	0.6077928	aldehyde dehydrogenase 4 family, member A1
814209 590774 969877 1469425 301122 199403	0.6064355 0.6055267 0.6041028	ESTs mitogen-activated protein kinase 13 synaptosomal-associated protein, 25kD SRY (sex determining region Y)-box 22 extracellular matrix protein 1 lectin, galactoside-binding, soluble, 8 (galectin 8)
813845 810331 770766 279970 490778	0.6014164 0.600234 0.5992634	RNA, U transporter 1 quiescin Q6 DKFZP564C1940 protein adenosine A2a receptor low molecular mass ubiquinone-binding protein (9.5kD)
288999 504959		small protein effector 1 of Cdc42 Homo sapiens mRNA; cDNA DKFZp586G0321 (from clone DKFZp586G0321)
42408 263727		hypothetical protein MGC4604 DNA segment, single copy probe LNS-CAI/LNS-CAII (deleted in polyposis
854763	0.5918867	Homo sapiens cDNA FLJ11341 fis, clone PLACE1010786
729975	0.5908615	meningioma expressed antigen 5 (hyaluronidase)
1733262 730313		BLu protein ESTs, Weakly similar to A43932 mucin 2 precursor, intestinal [H.sapiens]
1500000		H2B histone family, member B
1635681	0.5859113	retinoic acid induced 3 NADH dehydrogenase (ubiquinone) 1 alpha subcomplex, 2 (8kD, B8)
2325804	0.5840656	95 kDa retinoblastoma protein binding protein
2014856	0.5837779	major histocompatibility complex, class I-like sequence
	611443 782608 22778 128695 47853 814209 590774 969877 1469425 301122 199403 813845 810331 770766 279970 490778 288999 504959 42408 263727 854763 729975 1733262 730313 1500000 595037 1635681 2325804	6114430.61127137826080.6105794227780.61003221286950.608873478530.60779288142090.60769045907740.60719359698770.606435514694250.60410281994030.60271938138450.60189098103310.60141647707660.6002342799700.59926344907780.59574452889990.59556725049590.5953946424080.59302312637270.59263638547630.59188677299750.590861517332620.58918977303130.588757915000000.58717875950370.587070916356810.585911323258040.5840656

40173 796723		KIAA0807 protein Homo sapiens clone CDABP0014 mRNA sequence
239568 2018527		annexin A9 dolichyl-phosphate mannosyltransferase polypeptide 3
1649374	0.5731982	homogentisate 1,2-dioxygenase (homogentisate oxidase)
809357	0.571929	Bernardinelli-Seip congenital lipodystrophy 2 (seipin)
366132	0.5711651	succinate dehydrogenase complex, subunit C, integral membrane protein, 15kD
83358 1762111	0.5709886 0.5703701	ESTs natriuretic peptide receptor C/guanylate cyclase C (atrionatriuretic peptide receptor C)
244801 127646	0.5693836 0.5663395	Rho guanine exchange factor (GEF) 11
39884		IMP (inosine monophosphate) dehydrogenase 1
202577 588262		histamine N-methyltransferase Homo sapiens, Similar to RIKEN cDNA 2600001A11 gene, clone MGC:9907 IMAGE:3870073, mRNA, complete cds
813419	0.5602324	hydroxyacyl-Coenzyme A dehydrogenase, type II
1911343 859761		RAB26, member RAS oncogene family poliovirus receptor-related 2 (herpesvirus entry mediator B)
342181 1558108		B-cell CLL/lymphoma 2 ATP-binding cassette, sub-family C (CFTR/MRP), member 8
2017403 122394	0.5568495 0.5554851	regulator of G-protein signalling 3 guanine nucleotide binding protein (G protein), alpha 15 (Gq class)
824879 1708055 1456701 186301	0.554593 0.5545861	hypothetical protein MGC11275
126851 826622		hypothetical protein FLJ11160 KIAA0430 gene product

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151449	0.5503739	protein tyrosine phosphatase, non-receptor type 21
826668 76605 786072 2244196	0.5482232 0.5450231	KIAA0274 gene product nesca protein myosin IC B-cell receptor-associated protein BAP29
704414		small nuclear ribonucleoprotein polypeptides B and B1
739578	0.5402953	GPI-anchored metastasis-associated protein homolog
1404774 66599		parathyroid hormone-like hormone N-acetyltransferase 1 (arylamine N-acetyltransferase)
359250 488202		carbonic anhydrase IV ESTs, Weakly similar to YZ28_HUMAN HYPOTHETICAL PROTEIN ZAP128 [H.sapiens]
235986	0.5371486	wingless-type MMTV integration site family, member 11
214068 1500542 470092 589115	0.5341753 0.5319336	GATA-binding protein 3 regulator of G-protein signalling 11 like-glycosyltransferase matrix metalloproteinase 1 (interstitial collagenase)
810734 685516 171912	0.5304998	polymerase (DNA-directed), delta 4 putative G protein-coupled receptor Homo sapiens cDNA FLJ10960 fis, clone PLACE1000564
1858892 414992		hypothetical protein MGC4825 K562 cell-derived leucine-zipper-like protein 1
37708 713782	0.5283276 0.5273402	hypothetical protein MGC3101 a disintegrin and metalloproteinase domain 15 (metargidin)
471568	0.5255851	hematological and neurological expressed 1
812238 1631735	0.5237287 0.5228778	hypothetical protein MGC4692 Homo sapiens, clone IMAGE:3604336, mRNA, partial cds
74070 782497	0.5225238 0.5218113	endosulfine alpha Homo sapiens, clone IMAGE:3010666, mRNA, partial cds
502774	0.5204735	hypothetical protein FLJ20623

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1517171 50562 60565 1641894 51083	0.5195335 0.5192032 0.5187462	interleukin 2 receptor, alpha chromosome 8 open reading frame 4 lethal giant larvae (Drosophila) homolog 2 EST catenin (cadherin-associated protein), delta 2 (neural plakophilin-related arm-repeat protein)
296702 2018337		deiodinase, iodothyronine, type I glucosidase, beta; acid (includes glucosylceramidase)
1605426 1474149		hypothetical protein FLJ13352 poliovirus receptor-related 1 (herpesvirus entry mediator C; nectin)
26294 810017	0.5137265	RNB6 plasminogen activator, urokinase receptor
825822		hypothetical protein
810725	0.51329	ATPase, H+ transporting, lysosomal (vacuolar proton pump) 21kD
781019	0.5132896	paraoxonase 2
726658	0.5132765	non-metastatic cells 3, protein expressed in
1916575	0.5127478	BCL2-interacting killer (apoptosis-inducing)
1469148 726703		FGFR1 oncogene partner Homo sapiens clone 23736 mRNA sequence
725978	0.5108817	ESTs, Moderately similar to ALU7_HUMAN ALU SUBFAMILY SQ SEQUENCE CONTAMINATION WARNING ENTRY [H.sapiens]
1573946 51741 156363 177827 294397 826077 949938	0.5094912 0.5086284 0.5085582 0.505999 0.503983	programmed cell death 9 GTP-binding protein hypothetical protein FLJ12934 synaptotagmin VII DKFZP586A0522 protein pyruvate dehydrogenase (lipoamide) beta cystatin C (amyloid angiopathy and cerebral hemorrhage)
1957136 2018084 784140 810981	0.4999242 0.4984039	phenylethanolamine N-methyltransferase Ste-20 related kinase WD repeat domain 15 hypothetical protein FLJ20699

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627248 2302099 302031 183440 1609748 183200	0.4967769 0.49677 0.4967506 0.4955206	SBBI31 protein sialidase 3 (membrane sialidase) Ste20-related serine/threonine kinase arylsulfatase A hypothetical protein MGC10882 fumarylacetoacetate hydrolase (fumarylacetoacetase)
2306697 292770		neuromedin B Homo sapiens, clone IMAGE:3627860, mRNA, partial cds
856447 855563		interferon, gamma-inducible protein 30 v-erb-b2 avian erythroblastic leukemia viral oncogene homolog
839081	0.4939384	homolog of yeast long chain polyunsaturated fatty acid elongation enzyme 2
49273	0.4937997	solute carrier family 27 (fatty acid transporter), member 4
289505	0.4933344	ESTs, Moderately similar to S65657 alpha-1C-adrenergic receptor splice form 2 [H.sapiens]
782689	0.4930071	solute carrier family 6 (neurotransmitter transporter, creatine), member 8
767495	0.4922026	GLI-Kruppel family member GLI3 (Greig cephalopolysyndactyly
782537	0.490998	syndrome) Homo sapiens cDNA: FLJ22562 fis, clone HSI01814
855029 705274 202514	0.4904221	Ac-like transposable element diacylglycerol kinase, delta (130kD) DNA (cytosine-5-)-methyltransferase 3 alpha
725649	0.4878638	nuclear factor of activated T-cells, cytoplasmic, calcineurin-
1884404	0.4878341	dependent 4 KIAA0285 gene product
825296	0.4858551	low density lipoprotein receptor defect C complementing
839580 289857 491465 360778		ESTs phenylethanolamine N-methyltransferase hypothetical protein FLJ10035
	0.4814013	putative nuclear protein ORF1-FL49 ATPase, H+ transporting, lysosomal (vacuolar proton pump) 16kD
sd-71385		

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123614 1603583		hypothetical protein MGC4675 SH3 domain binding glutamic acid-rich protein like
782688	0.4780253	dynein, axonemal, light intermediate polypeptide
784150 770869 455269 365358 795256 756666	0.4762758 0.4761389 0.4756564	RAB31, member RAS oncogene family hypothetical protein MGC2592 pM5 protein NPD007 protein protein phosphatase 1, catalytic subunit, alpha isoform
767706 488505 344073		
1558151	0.4723511	basic leucine zipper transcription factor, ATF-like
823615	0.471739	Homo sapiens cDNA: FLJ21245 fis, clone COL01184
49351 1541711	0.4697521 0.4694413	SEX gene Homo sapiens cDNA: FLJ21513 fis, clone COL05778
868652 430318 143426 951125 66535 813281 726699	0.4681908 0.4679928 0.4675003 0.466705 0.4659334	complement component 4B parvalbumin ras homolog gene family, member B peroxisomal D3,D2-enoyl-CoA isomerase ornithine decarboxylase antizyme 2 WW domain-containing protein 1 ESTs, Weakly similar to AAB47496 NG5 [H.sapiens]
1492468 1492426 489106 488422 278430 1518890	0.4635224 0.4628623 0.4626549 0.4612914	DEME-6 protein chromosome 19 open reading frame 3 hypothetical protein FLJ11210 high-mobility group 20B kinesin family member 5C metallothionein-like 5, testis-specific (tesmin)
1557637	0.4607567 0.4588694	homolog of yeast MOG1 seven transmembrane protein TM7SF3 ESTs trefoil factor 3 (intestinal)

810402 767761 1473289	0.4565666	hypothetical protein DKFZP434B168 protein protective protein for beta-galactosidase (galactosialidosis)
773142 1652310	0.4530754 0.4525789	hypothetical protein MGC2827 Homo sapiens, clone MGC:19613 IMAGE:3833049, mRNA, complete cds
812143	0.4519186	fibronectin leucine rich transmembrane protein 3
2326057 1075635 825327	0.4502629	MLN51 protein MLSN1- and TRP-related Homo sapiens cDNA FLJ14105 fis, clone MAMMA1001202
809838 786545	0.4466933 0.4453714	protein phosphatase 1, regulatory (inhibitor) subunit 3B
183556	0.4452493	gap junction protein, alpha 4, 37kD (connexin 37)
503889 725321 731459	0.44288	hypothetical protein FLJ10656 estrogen receptor 1 Homo sapiens cDNA: FLJ22296 fis, clone HRC04468
752631	0.4426961	fibroblast growth factor receptor 3 (achondroplasia, thanatophoric dwarfism)
1558642		hypothetical protein MGC2771
120271		hypothetical protein MGC4692
487932		synaptotagmin-like 2
1592715		Homer, neuronal immediate early gene, 3
308466 840882		GTP-binding protein Sara nucleotide binding protein
823727		Homo sapiens, clone IMAGE:2905978, mRNA, partial cds
1636092 1505038 490965	0.4404649 0.43988 0.4389746	· · · · · · · · · · · · · · · · · · ·
346696	0.437234	TEA domain family member 4
345032	0.4360992	ESTs
487733 810612	0.4356439 0.4352602	S100 calcium-binding protein A11 (calgizzarin)
122077 417801	0.434495 0.4325892	putative membrane protein mitochondrial ribosomal protein L27

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878406 1947381 232789		metaxin 1 hypothetical protein FLJ22329 polymerase (RNA) II (DNA directed) polypeptide J (13.3kD)
809944 2019101		KIAA0310 gene product angiotensinogen (serine (or cysteine) proteinase inhibitor, clade A (alpha-1 antiproteinase, antitrypsin), member 8)
1693357 75859	0.4284863 0.4274493 -1.6337267 -1.4515145	coagulation factor XII (Hageman factor) mitochondrial intermediate peptidase endothelin 2 N-myc downstream-regulated gene 2 SRY (sex determining region Y)-box 10 Homo sapiens cDNA: FLJ22528 fis, clone HRC12825
810002 712139 815737	-1.2371074	Homo sapiens, clone MGC:19762 IMAGE:3636045, mRNA, complete cds ADP-ribosylation factor-like 7 ATP synthase, H+ transporting, mitochondrial F1 complex, alpha subunit, isoform 1, cardiac muscle
503671	-1.1528584	Homo sapiens cDNA FLJ14368 fis, clone HEMBA1001122
345670	-1.135654	ESTs, Moderately similar to I59348 CCAAT binding transcription factor CBF subunit C - rat [R.norvegicus]
753038 741139 839736 2013178 1577736 788234	-1.1025542 -1.1019142 -1.0884449 -1.076549 -1.0610946	kinesin family member C3 eyes absent (Drosophila) homolog 2 crystallin, alpha B death associated transcription factor 1 epidermal growth factor (beta-urogastrone) inhibitor of DNA binding 4, dominant negative helix-loop-helix protein UDP-GlcNAc:betaGal beta-1,3-N-acetylglucosaminyltransferase
		3
757873	-1.0188169	cyclin-dependent kinase 5, regulatory subunit 1 (p35)
291478 486683		runt-related transcription factor 3 Homo sapiens mRNA; cDNA DKFZp564J0323 (from clone DKFZp564J0323)
290378 772913 sd-71385	-1.0043647 -0.9823487	podocalyxin-like calreticulin

P A T E N T Atty Dkt: 485772004300

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544639 814798	-0.9767278 -0.9611553	ESTs aldehyde dehydrogenase 1 family, member A3
131839 840266		folate receptor 1 (adult) Homo sapiens cDNA: FLJ22667 fis, clone HSI08385
1555924 756708		CSR1 protein potassium intermediate/small conductance calcium-activated channel, subfamily N, member 4
838478 742562 1469377 884462 796542 32493 284592 788136	-0.9269494 -0.9086679 -0.9065675 -0.9052403 -0.9043423 -0.9019931	neurocalcin delta chromosome 16 open reading frame 5 lipoma HMGIC fusion partner-like 2 Down syndrome critical region gene 1 ets variant gene 5 (ets-related molecule) integrin, alpha 6 PRO1659 protein phosphodiesterase 4B, cAMP-specific (dunce (Drosophila)-homolog phosphodiesterase E4)
1161564 1635320	-0.8865787 -0.8845852	desmuslin amiloride-sensitive cation channel 2, neuronal
344720 1534700 193913	-0.8804685	glycophorin C (Gerbich blood group) KIAA0830 protein v-yes-1 Yamaguchi sarcoma viral related oncogene homolog
71087	-0.870641	v-maf musculoaponeurotic fibrosarcoma (avian) oncogene family, protein F
300632	-0.8586688	hypothetical protein FLJ21044 similar to Rbig1
416676 69002		pellino (Drosophila) homolog 1 PPAR(gamma) angiopoietin related protein
742685	-0.852598	disabled (Drosophila) homolog 2 (mitogen-responsive
140574	-0.8498606	phosphoprotein) small inducible cytokine subfamily D (Cys-X3-Cys), member 1 (fractalkine, neurotactin)
23831 470393		aldolase C, fructose-bisphosphate matrix metalloproteinase 7 (matrilysin, uterine)
1474337	-0.844539	phosphorylase, glycogen; brain
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753301	-0.8393651	carcinoembryonic antigen-related cell adhesion molecule 1 (biliary glycoprotein)
51078	_n 8348835	ribosomal protein L44
3172883		ESTs, Weakly similar to 1709359A dopamine D4 receptor [H.sapiens]
877621	-0.8289253	nGAP-like protein
811920		interleukin 11 receptor, alpha
156211		ATPase, H+ transporting, lysosomal (vacuolar proton pump), beta polypeptide, 56/58kD, isoform 1 (Renal tubular acidosis with deafness)
760299	-0.8251086	dickkopf (Xenopus laevis) homolog 3
285377		pellino (Drosophila) homolog 2
345034		small inducible cytokine subfamily B (Cys-X-Cys), member 14 (BRAK)
298122	-0.8158372	frizzled (Drosophila) homolog 7
842896	-0.8123068	hypothetical protein DKFZp762L0311
843070		nucleoporin 88kD
1257131	-0.8069059	·
796539	-0.803124	KRAB-associated protein 1
139660	-0.79833	ESTs
666879	-0.7950485	annexin A8
200814	-0.7920988	membrane metallo-endopeptidase (neutral endopeptidase, enkephalinase, CALLA, CD10)
2016775	-0.7914834	G protein-coupled receptor, family C, group 5, member B
1946448	-0.790849	caveolin 2
1473471	-0.7906678	KIAA0194 protein
1556433	-0.7877792	GRO3 oncogene
1636156	-0.7827089	hypothetical protein FLJ21709
40299	-0.78007	growth differentiation factor 10
1636166		KIAA0668 protein
1609665		BarH-like homeobox 2
250797		hypothetical protein FLJ20038
220395	-0.7687195	hypothetical protein FLJ23293 similar to ARL-6 interacting protein-2
208718		annexin A1
41869		hypothetical protein FLJ11017
490023		hypothetical protein MGC2648
1607229		tumor protein D52-like 1
133236	-0.7561129	RNA binding motif protein, X chromosome
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	785733 32489 70201 611481 1554167 768571 783698 341759	-0.7554682 hypothetical protein FLJ12892 -0.7521767 hypothetical protein DKFZp566A1524 -0.7513314 mitochondrial solute carrier -0.7507264 HMG-box transcription factor TCF-3 -0.7503289 hypothetical protein FLJ14529 -0.7476813 SRY (sex determining region Y)-box 8 -0.7411614 lipin 1 -0.7393558 lung type-I cell membrane-associated glycoprotein
भूतमा देशमी भित्य भू	665384 142259	-0.7388404 KIAA1609 protein -0.7380151 tumor necrosis factor alpha-inducible cellular protein containing leucine zipper domains; Huntingtin interacting protein L; transcrption factor IIIA-interacting protein
	810057 27544 739193 813256	-0.7363941 vasoactive intestinal peptide receptor 1 -0.7342174 prominin (mouse)-like 1 -0.7338774 cellular retinoic acid-binding protein 1 -0.7334866 ATP-binding cassette, sub-family B (MDR/TAP), member 1
ा भेगा भेगा भेगा भेगा भेगा	293964 272706 209537 67741 823714	-0.7275302 butyrophilin, subfamily 3, member A1 -0.7274769 alpha2,3-sialyltransferase -0.7245778 zinc finger protein 221 -0.7229251 PP2135 protein -0.7215045 nuclear receptor co-repressor/HDAC3 complex subunit
	471196 813265	-0.7192752 integral membrane protein 3 -0.7192289 Homo sapiens mRNA; cDNA DKFZp564H1916 (from clone DKFZp564H1916)
	42681 1601845 725152 22917	-0.718704 NY-REN-25 antigen -0.7178673 Ca2+-promoted Ras inactivator -0.7173869 hypothetical protein DKFZp762A227 -0.7173328 Homo sapiens mRNA; cDNA DKFZp761M0111 (from clone DKFZp761M0111)
	703541 529843	-0.7118124 KIAA1858 protein -0.7117107 ESTs, Moderately similar to JC5238 galactosylceramide-like protein, GCP [H.sapiens]
	377461 1569187	-0.7093974 caveolin 1, caveolae protein, 22kD -0.7091259 heparan sulfate (glucosamine) 3-O-sulfotransferase 4

1456118	-0.7074166	proteasome (prosome, macropain) subunit, beta type, 9 (large multifunctional protease 2)
289760	-0.7058375	ESTs, Highly similar to T00391 hypothetical protein KIAA0612 [H.sapiens]
341774	-0.7044768	major histocompatibility complex, class I, A
75078 505864 415191 188335	-0.700303	ESTs RalGDS-like gene KIAA0161 gene product egf-like module containing, mucin-like, hormone receptor-like sequence 2
132711 1864302		Kruppel-like factor 5 (intestinal) E74-like factor 5 (ets domain transcription factor)
344854 454970 79629		ankyrin repeat domain 3 DKFZP434G032 protein
767068 378813		DKFZP586G1517 protein secretory leukocyte protease inhibitor (antileukoproteinase)
725076 491403		5'-nucleotidase (purine), cytosolic type B tumor necrosis factor receptor superfamily, member 1B
43764	-0.6743049	hypothetical protein FLJ14033 similar to hypoxia inducible factor 3, alpha subunit
725622 431231	-0.6731681 -0.6692022	•
488956 72778	-0.6682156 -0.6614808	CUG triplet repeat, RNA-binding protein 2 caspase 7, apoptosis-related cysteine protease
360254 76182 773487 162308 283023	-0.6603322 -0.6598647 -0.6580605 -0.6577611	chemokine (C-X3-C) receptor 1
196992 303109	-0.6571713	aldo-keto reductase family 1, member C1 (dihydrodiol dehydrogenase 1; 20-alpha (3-alpha)-hydroxysteroid dehydrogenase) purinergic receptor (family A group 5)
897731	-0.6548064	latrophilin

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289428	-0.6542789	neurotrophic tyrosine kinase, receptor, type 2
196435 504791 812975 488404	-0.6527728	ESTs glutathione S-transferase A4 KIAA0172 protein Homo sapiens clone TUA8 Cri-du-chat region mRNA
345056 786069 48518	-0.6484842	KIAA1404 protein beta-site APP-cleaving enzyme ATP-binding cassette, sub-family A (ABC1), member 5
812048	-0.6438062	prion protein (p27-30) (Creutzfeld-Jakob disease, Gerstmann-Strausler-Scheinker syndrome, fatal familial insomnia)
868396	-0.6426678	tyrosine 3-monooxygenase/tryptophan 5-monooxygenase activation protein, eta polypeptide
70245	-0.6422928	Homo sapiens mRNA full length insert cDNA clone EUROIMAGE 50374
40027	-0.6418674	
840944		early growth response 1
809784		kallikrein 6 (neurosin, zyme)
1881774	-0.6332545	
52419		Friedreich ataxia region gene X123
120106		caspase 1, apoptosis-related cysteine protease (interleukin 1, beta, convertase)
725680	-0.6295778	transcription factor AP-2 gamma (activating enhancer-binding protein 2 gamma)
1493218	-0.6292715	hypothetical protein FLJ22297
2018423		death-associated protein kinase 2
1486082		heparin-binding growth factor binding protein
868169	-0.6224058	lipoprotein lipase
209137		gamma-aminobutyric acid (GABA) A receptor, epsilon
		ga
248886	-0.6161134	rab3 GTPase-activating protein, non-catalytic subunit (150kD)
73252	-0.6115602	3-hydroxy-3-methylglutaryl-Coenzyme A synthase 1 (soluble)
612274 346545 84464	-0.6098367	tubulin, alpha 1 (testis specific) laminin, beta 1 hypothetical protein FLJ12806
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	-0.6078858	forkhead box O1A (rhabdomyosarcoma) chromosome 2 open reading frame 2 Homo sapiens mRNA for FLJ00074 protein, partial cds
796904 418159 323238	-0.6038708	pleiomorphic adenoma gene-like 1 synaptogyrin 1 GRO1 oncogene (melanoma growth stimulating activity, alpha)
1570427 1500815	-0.601608 -0.6008126	hypothetical protein MGC4309 Homo sapiens cDNA: FLJ22130 fis, clone HEP19632
416959 782460		nuclear factor I/B Homo sapiens cDNA FLJ10500 fis, clone NT2RP2000369
824602 878798 32299 277571 703964 796498 504927	-0.5956349 -0.5945347 -0.59335 -0.5918604 -0.5900328	interferon, gamma-inducible protein 16 beta-2-microglobulin inositol(myo)-1(or 4)-monophosphatase 2 KIAA1706 protein inositol polyphosphate phosphatase-like 1 hypothetical protein FLJ14007 epithelial protein up-regulated in carcinoma, membrane associated protein 17
813520 341763	-0.5889772 -0.5868818	EphB3 caspase 5, apoptosis-related cysteine protease
1856063 825356		tweety (Drosophila) homolog 1 Homo sapiens cDNA FLJ11997 fis, clone HEMBB1001458
745011	-0.5836864	Homo sapiens mRNA for KIAA1750 protein, partial cds
812256	-0.5832398	Homo sapiens cDNA: FLJ21693 fis, clone COL09609
		PP1201 protein granzyme A (granzyme 1, cytotoxic T-lymphocyte-associated serine esterase 3)
815142	-0.5782827	spastic ataxia of Charlevoix-Saguenay (sacsin)
80727	-0.5766546	receptor tyrosine kinase-like orphan receptor 1
45099	-0.5753698	regucalcin (senescence marker protein-30)
415816	-0.5750184	ESTs
od 71205		

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306798	-0.5742664	NGFI-A binding protein 1 (EGR1 binding protein 1)
740620	-0.5735888 -0.5730915 -0.5729078	dendritic cell protein ESTs tropomyosin 2 (beta) lymphocyte antigen 6 complex, locus E peroxisomal membrane protein 3 (35kD, Zellweger syndrome)
37671	-0.5712258	checkpoint suppressor 1 hypothetical protein FLJ21610 v-akt murine thymoma viral oncogene homolog 2
132637	-0.5694152	grancalcin, EF-hand calcium-binding protein
34093 1881689 1883327	-0.5684295	EST hypothetical protein FLJ20281
781017	-0.5676944	early growth response 2 (Krox-20 (Drosophila) homolog)
	-0.5673709 -0.5667198	ESTs Homo sapiens mRNA; cDNA DKFZp434G0972 (from clone DKFZp434G0972)
470128 1880885 811848	-0.5652125 -0.5638973 -0.5637994 -0.563564	
1911663 66491	-0.5611985 -0.5610314 -0.5579872	
1626996	-0.5573507	c-fos induced growth factor (vascular endothelial growth factor D)

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	-0.5558155	ESTs four and a half LIM domains 3 heat shock 27kD protein 2 hypothetical protein FLJ22167
		lactate dehydrogenase C
809998		amylase, alpha 2A; pancreatic
1734754	-0.5512835	hypothetical protein B
27769	-0.5501268	
950355	-0.550114	ESTs, Weakly similar to S13495 pregnancy zone protein [H.sapiens]
144924	-0.5498123	chromosome 6 open reading frame 5
502436	-0.5489864	insulin receptor substrate 2
138672	-0.5483104	
160192	-0.5482009	ESTs, Weakly similar to 2004399A chromosomal protein [H.sapiens]
768007	-0.547374	hypothetical protein
130201		intercellular adhesion molecule 2
1587847	-0.5466442	minichromosome maintenance deficient (mis5, S. pombe) 6
811088	-0.5449844 -0.5445374 -0.5440514	hypothetical protein FLJ21841 ephrin-B3
530958 810097	-0.5423532 -0.5423072	smoothened (Drosophila) homolog Homo sapiens cDNA: FLJ21721 fis, clone COLF0381
207735	-0.5422739	serine (or cysteine) proteinase inhibitor, clade A (alpha-1 antiproteinase, antitrypsin), member 1
70054		
79254 85634		MHC class I region ORF complement component 1, s subcomponent

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970590	-0.5383645	Homo sapiens mRNA; cDNA DKFZp434A115 (from clone DKFZp434A115)
		KIAA0603 gene product Homo sapiens cDNA: FLJ21028 fis, clone CAE07155
153760 247616	-0.5354869	EphB1 Iipoma HMGIC fusion partner
1686766 2009491 611532 811149 844703	-0.5334821 -0.5320107 -0.5319742	Rag D protein cyclin-E binding protein 1 troponin I, skeletal, fast chromosome 9 open reading frame 3 Sam68-like phosphotyrosine protein, T-STAR
70749	-0.5295091	Homo sapiens cDNA: FLJ21874 fis, clone HEP02488
324927 491519		KIAA0375 gene product Homo sapiens clone 24775 mRNA sequence
754040 767765		NS1-associated protein 1 GTP-binding protein overexpressed in skeletal muscle
2028722 509458 360518	-0.5266016	osteoblast specific factor 2 (fasciclin I-like) hypothetical protein from clone 643 guanine nucleotide binding protein (G protein), gamma transducing activity polypeptide 1
40021	-0.5240809	hairless protein (putative single zinc finger transcription factor protein, responsible for autosomal recessive universal congenital alopecia, HR gene)
1901310 33949		KIAA1209 protein phosphoribosyl pyrophosphate synthetase-associated protein 1
814316 488130		ribosomal protein L13 Homo sapiens cDNA FLJ20767 fis, clone COL06986
462939	-0.5200097	ESTs

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770848	-0.5188717	ESTs, Weakly similar to ALU1_HUMAN ALU SUBFAMILY J SEQUENCE CONTAMINATION WARNING ENTRY [H.sapiens]
245296 753034	-0.5184238 -0.5182019	RAD52 (S. cerevisiae) homolog nuclear factor I/X (CCAAT-binding transcription factor)
712829 1577920 814526 773301 77972 1704155 1572298	-0.5161377 -0.5156162 -0.5147295 -0.5144916 -0.5133015	
		retinoic acid receptor, gamma DnaJ (Hsp40) homolog, subfamily A, member 2
712401	-0.5107426	phosphoinositide-3-kinase, catalytic, delta polypeptide
713839	-0.5088558	transcription factor AP-4 (activating enhancer-binding protein 4)
180803	-0.5083354	inositol polyphosphate-1-phosphatase

Example V: Genes for discriminating between normal and DCIS

-0.5075293 hypothetical protein MGC3232

As shown in Table 4 below, 1300 genes were identified as being able to discriminate between normal and DCIS.

Table 4

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ClonelD	Weight	Description
1323448	1.397391	cysteine-rich protein 1 (intestinal)
788654	1.2615322	growth factor receptor-bound protein 2
1505038	1.1904802	hypothetical protein FLJ20171
745606	1.1825585	hypothetical protein PP591

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565319	1.0969429	Homo sapiens mRNA; cDNA DKFZp564B1264 (from clone DKFZp564B1264)
35147	1.0886084	ESTs, Weakly similar to unnamed protein product [H.sapiens]
178805	1.0274897	Human DNA sequence from clone RP5-850E9 on chromosome 20. Contains part of the gene for a novel C2H2 type zinc finger protein similar to Drosophila Scratch (Scrt), Slug and Xenopus Snail, a novel gene similar to Drosophila CG6762, STSs, GSSs and five CpG
796469	1.0269115	HSPC150 protein similar to ubiquitin-conjugating enzyme
1609836	1.0164283	glutamate-ammonia ligase (glutamine synthase)
366132	1.013995	succinate dehydrogenase complex, subunit C, integral membrane protein, 15kD
1500000 812238 595037 1554549 488964 138189 283919 823598	1.0043787 0.990461 0.9764206 0.9569176 0.9386802 0.937383 0.9153521	H2B histone family, member B hypothetical protein MGC4692 retinoic acid induced 3 hydroxyacyl glutathione hydrolase H2A histone family, member O Wolfram syndrome 1 (wolframin) H2A histone family, member L proteasome (prosome, macropain) 26S subunit, non-ATPase, 12 stearoyl-CoA desaturase (delta-9-desaturase)
503215 811774 122077	0.9108688 0.9073627 0.9021901 0.9019233	RAB26, member RAS oncogene family pilin-like transcription factor CGI-49 protein putative membrane protein HSPC003 protein Ric (Drosophila)-like, expressed in many tissues
685516 359887 131094	0.8692353	putative G protein-coupled receptor translocase of inner mitochondrial membrane 17 (yeast) homolog A Homo sapiens cDNA: FLJ21587 fis, clone COL06946
1492463	0.8634521	selenoprotein X, 1

	0	1	
_	0	1	•

843195 796723		phosphoserine phosphatase Homo sapiens clone CDABP0014 mRNA sequence
488202	0.8463961	ESTs, Weakly similar to YZ28_HUMAN HYPOTHETICAL PROTEIN ZAP128 [H.sapiens]
280375 290841 1917941	0.8392836 0.8346933 0.8340565 0.8296141	chromosome 19 open reading frame 3 PRO2000 protein H2B histone family, member A purine-rich element binding protein B SRY (sex determining region Y)-box 22 ESTs, Weakly similar to CA13_HUMAN COLLAGEN ALPHA 1(III) CHAIN PRECURSOR [H.sapiens]
471568	0.8264546	hematological and neurological expressed 1
		hypothetical protein FLJ13187 ESTs, Weakly similar to N-WASP [H.sapiens]
731044 244801 2054635	0.8066357	glutaredoxin 2 Rho guanine exchange factor (GEF) 11 proteasome (prosome, macropain) subunit, alpha type, 7
1605426 1709791 470061 1640821	0.7837549 0.77977	hypothetical protein FLJ13352 BAI1-associated protein 1 seven in absentia (Drosophila) homolog 2 ESTs, Weakly similar to I78885 serine/threonine-specific protein kinase [H.sapiens]
199403	0.7754748	lectin, galactoside-binding, soluble, 8 (galectin 8)
		glucose phosphate isomerase antigen identified by monoclonal antibodies 12E7, F21 and O13
239568 841679		annexin A9 calcium and integrin binding protein (DNA-dependent protein kinase interacting protein)
729975	0.7649717	meningioma expressed antigen 5 (hyaluronidase)
2016648	0.7633048	Homo sapiens mRNA; cDNA DKFZp434N1728 (from clone DKFZp434N1728)
2052113 sd-71385	0.7627076	hypothetical protein FLJ10903

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347373	0.7563599	transcription elongation factor B (SIII), polypeptide 1 (15kD, elongin C)
814054	0.7539407	KIAA0040 gene product
741977		B-factor, properdin
272529		phosphomannomutase 2
824879	0.7432221	
122241	0.7394284	proteasome (prosome, macropain) subunit, beta type, 2
744417		carnitine acetyltransferase
868128		JM4 protein
241348		prenylcysteine lyase
509823	0.7304507	carcinoembryonic antigen-related cell adhesion molecule 6
		(non-specific cross reacting antigen)
1636092	0.7270017	hypothetical protein FLJ20657
42408	0.725109	
686552	0.7231319	golgi phosphoprotein 1
898032	0.7216	KIAA0097 gene product
754628	0.7195265	
826256	0.7190206	transmembrane 7 superfamily member 1 (upregulated in
		kidney)
2043167		BCL2-associated athanogene 3
827171	0.7154964	
502774		hypothetical protein FLJ20623
781097		reticulon 3
810725	0.7115051	ATPase, H+ transporting, lysosomal (vacuolar proton pump)
		21kD
768570		hypothetical protein FLJ11280
813281		WW domain-containing protein 1
39884	0.7075527	IMP (inosine monophosphate) dehydrogenase 1
969877	0.707469	synaptosomal-associated protein, 25kD
704254	0.7061648	a disintegrin and metalloproteinase domain 8
000004	0.7000000	
236034	0.7039329	uncoupling protein 2 (mitochondrial, proton carrier)
782428	0.701077	KIAA0250 gene product
489351	0.7003616	
37708	0.6986007	
271472		C3HC4-like zinc finger protein
782608	0.6978285	mitochondrial ribosomal protein L9

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704414	0.6940045	small nuclear ribonucleoprotein polypeptides B and B1
288999 51773 124447 470099	0.6925666 0.6918546	small protein effector 1 of Cdc42 hypothetical protein MGC3077 KIAA1184 protein HT002 protein; hypertension-related calcium-regulated gene
429799	0.6915155	
1858892 796694		gene 2 hypothetical protein MGC4825 baculoviral IAP repeat-containing 5 (survivin)
284261	0.6890531	uncharacterized hematopoietic stem/progenitor cells protein MDS030
814528	0.6878182	Homo sapiens cDNA: FLJ22139 fis, clone HEP20959
788654 68636 308466 1574058	0.6800103	hypothetical protein MGC2477 GTP-binding protein Sara 1-acylglycerol-3-phosphate O-acyltransferase 2 (lysophosphatidic acid acyltransferase, beta)
79520 1473289	0.6769129 0.674396	RAB2, member RAS oncogene family protective protein for beta-galactosidase (galactosialidosis)
365738 768064	0.6740773 0.673441	ESTs cytochrome P450, subfamily I (aromatic compound-inducible), polypeptide 1
773922	0.6730903	KIAA0005 gene product
1869201	0.6727191	71
686172	0.6724684	
810558	0.6698194	proteasome (prosome, macropain) 26S subunit, ATPase, 4
753299	0.6694573	hypothetical protein FLJ10504
725454		CDC28 protein kinase 2
470124		RAD1 (S. pombe) homolog
2014034	0.6620303	methylene tetrahydrofolate dehydrogenase (NAD+ dependent), methenyltetrahydrofolate cyclohydrolase
209066 859761	0.6611038 0.6573769	poliovirus receptor-related 2 (herpesvirus entry mediator B)
589232 sd-71385	0.6551447	hypothetical protein FLJ11506

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340558	0.6550804	actin related protein 2/3 complex, subunit 5 (16 kD)
186768	0.6550741	Homo sapiens, clone IMAGE:3604869, mRNA
813629 1492780		YME1 (S.cerevisiae)-like 1 Homo sapiens cDNA FLJ14459 fis, clone HEMBB1002409
809944 83363		KIAA0310 gene product protein-L-isoaspartate (D-aspartate) O-methyltransferase
149355	0.6520333	translocating chain-associating membrane protein
752631	0.650399	fibroblast growth factor receptor 3 (achondroplasia, thanatophoric dwarfism)
785616	0.6484744	·
76605	0.6484522	nesca protein
742707	0.6476434	ESTs, Weakly similar to MUC2_HUMAN MUCIN 2 PRECURSOR [H.sapiens]
325606	0.647638	hypothetical protein MGC14353
769921	0.6464245	ubiquitin carrier protein E2-C
1435003	0.6457486	tumor suppressing subtransferable candidate 1
1473922	0.6450042	actin related protein 2/3 complex, subunit 3 (21 kD)
1170022	0.0100012	
		N-acetylneuraminic acid phosphate synthase; sialic acid
1456348		N-acetylneuraminic acid phosphate synthase; sialic acid synthase
1456348	0.6417276 0.6416738	N-acetylneuraminic acid phosphate synthase; sialic acid synthase
1456348 1500162	0.6417276 0.6416738 0.6410435	N-acetylneuraminic acid phosphate synthase; sialic acid synthase ESTs
1456348 1500162 1573251	0.6417276 0.6416738 0.6410435 0.640944	N-acetylneuraminic acid phosphate synthase; sialic acid synthase ESTs peroxisomal long-chain acyl-coA thioesterase ADP-ribosyltransferase (NAD+; poly (ADP-ribose) polymerase)
1456348 1500162 1573251 46248 49351	0.6417276 0.6416738 0.6410435 0.640944 0.6386132	N-acetylneuraminic acid phosphate synthase; sialic acid synthase ESTs peroxisomal long-chain acyl-coA thioesterase ADP-ribosyltransferase (NAD+; poly (ADP-ribose) polymerase) SEX gene
1456348 1500162 1573251 46248	0.6417276 0.6416738 0.6410435 0.640944 0.6386132 0.6375787	N-acetylneuraminic acid phosphate synthase; sialic acid synthase ESTs peroxisomal long-chain acyl-coA thioesterase ADP-ribosyltransferase (NAD+; poly (ADP-ribose) polymerase)
1456348 1500162 1573251 46248 49351 824052 1420370	0.6417276 0.6416738 0.6410435 0.640944 0.6386132 0.6375787 0.6344209	N-acetylneuraminic acid phosphate synthase; sialic acid synthase ESTs peroxisomal long-chain acyl-coA thioesterase ADP-ribosyltransferase (NAD+; poly (ADP-ribose) polymerase) SEX gene chromosome 6 open reading frame 1 biliverdin reductase B (flavin reductase (NADPH))
1456348 1500162 1573251 46248 49351 824052 1420370 842994	0.6417276 0.6416738 0.6410435 0.640944 0.6386132 0.6375787 0.6344209 0.6343858	N-acetylneuraminic acid phosphate synthase; sialic acid synthase ESTs peroxisomal long-chain acyl-coA thioesterase ADP-ribosyltransferase (NAD+; poly (ADP-ribose) polymerase) SEX gene chromosome 6 open reading frame 1 biliverdin reductase B (flavin reductase (NADPH)) cathepsin Z
1456348 1500162 1573251 46248 49351 824052 1420370	0.6417276 0.6416738 0.6410435 0.640944 0.6386132 0.6375787 0.6344209 0.6343858	N-acetylneuraminic acid phosphate synthase; sialic acid synthase ESTs peroxisomal long-chain acyl-coA thioesterase ADP-ribosyltransferase (NAD+; poly (ADP-ribose) polymerase) SEX gene chromosome 6 open reading frame 1 biliverdin reductase B (flavin reductase (NADPH))
1456348 1500162 1573251 46248 49351 824052 1420370 842994	0.6417276 0.6416738 0.6410435 0.640944 0.6386132 0.6375787 0.6344209 0.6343858 0.6341098 0.633308	N-acetylneuraminic acid phosphate synthase; sialic acid synthase ESTs peroxisomal long-chain acyl-coA thioesterase ADP-ribosyltransferase (NAD+; poly (ADP-ribose) polymerase) SEX gene chromosome 6 open reading frame 1 biliverdin reductase B (flavin reductase (NADPH)) cathepsin Z S100 calcium-binding protein A11 (calgizzarin) hypothetical protein MGC4675
1456348 1500162 1573251 46248 49351 824052 1420370 842994 810612 123614 124781	0.6417276 0.6416738 0.6410435 0.640944 0.6386132 0.6375787 0.6344209 0.6343858 0.6341098 0.633308 0.6324418	N-acetylneuraminic acid phosphate synthase; sialic acid synthase ESTs peroxisomal long-chain acyl-coA thioesterase ADP-ribosyltransferase (NAD+; poly (ADP-ribose) polymerase) SEX gene chromosome 6 open reading frame 1 billiverdin reductase B (flavin reductase (NADPH)) cathepsin Z S100 calcium-binding protein A11 (calgizzarin) hypothetical protein MGC4675 squalene epoxidase
1456348 1500162 1573251 46248 49351 824052 1420370 842994 810612 123614 124781 814378	0.6417276 0.6416738 0.6410435 0.640944 0.6386132 0.6375787 0.6344209 0.6343858 0.6341098 0.633308 0.6324418 0.6317382	N-acetylneuraminic acid phosphate synthase; sialic acid synthase ESTs peroxisomal long-chain acyl-coA thioesterase ADP-ribosyltransferase (NAD+; poly (ADP-ribose) polymerase) SEX gene chromosome 6 open reading frame 1 biliverdin reductase B (flavin reductase (NADPH)) cathepsin Z S100 calcium-binding protein A11 (calgizzarin) hypothetical protein MGC4675 squalene epoxidase serine protease inhibitor, Kunitz type, 2
1456348 1500162 1573251 46248 49351 824052 1420370 842994 810612 123614 124781 814378	0.6417276 0.6416738 0.6410435 0.640944 0.6386132 0.6375787 0.6344209 0.6343858 0.6341098 0.633308 0.6324418 0.6317382	N-acetylneuraminic acid phosphate synthase; sialic acid synthase ESTs peroxisomal long-chain acyl-coA thioesterase ADP-ribosyltransferase (NAD+; poly (ADP-ribose) polymerase) SEX gene chromosome 6 open reading frame 1 billiverdin reductase B (flavin reductase (NADPH)) cathepsin Z S100 calcium-binding protein A11 (calgizzarin) hypothetical protein MGC4675 squalene epoxidase

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289978 67765 742595 727078	0.6300886 0.6265846	ubiquitin-like 4 carboxypeptidase M cyclin-dependent kinase 5 Homo sapiens cDNA: FLJ23602 fis, clone LNG15735
487733 366067	0.6262778 0.625758	cerebellar degeneration-related protein (62kD)
292770	0.6231261	Homo sapiens, clone IMAGE:3627860, mRNA, partial cds
366353 810063	0.6203012 0.620086	DKFZP564C186 protein growth factor, erv1 (S. cerevisiae)-like (augmenter of liver regeneration)
810124	0.6162712	platelet-activating factor acetylhydrolase, isoform lb, gamma subunit (29kD)
713782	0.6149877	a disintegrin and metalloproteinase domain 15 (metargidin)
782608 530197	0.6137782 0.6122721	ESTs, Moderately similar to ALU8_HUMAN ALU SUBFAMILY SX SEQUENCE CONTAMINATION WARNING ENTRY [H.sapiens]
44292	0.6122043	Homo sapiens mRNA; cDNA DKFZp434C107 (from clone DKFZp434C107)
	0.6104002 0.6102239	Ste-20 related kinase fibulin 2 hydroxysteroid (17-beta) dehydrogenase 7 Homo sapiens cDNA FLJ14105 fis, clone MAMMA1001202
842980	0.6081528	developmentally regulated GTP-binding protein 1
811761 1601947	0.6045809 0.6045149	Nijmegen breakage syndrome 1 (nibrin) cytochrome c oxidase subunit VIIa polypeptide 2 (liver)
1474955	0.6045103	TATA box binding protein (TBP)-associated factor, RNA polymerase II, N, 68kD (RNA-binding protein 56)
1758590	0.6040962	fatty-acid-Coenzyme A ligase, long-chain 3
1456701 207288 1631699	0.603346	B-cell CLL/lymphoma 9 insulin induced gene 1 valosin-containing protein

1635681	0.6022549	NADH dehydrogenase (ubiquinone) 1 alpha subcomplex, 2 (8kD, B8)
768452	0.6018854	Homo sapiens EST from clone 491476, full insert
897806	0.6015789	hypoxia-inducible factor 1, alpha subunit (basic helix-loop-helix transcription factor)
108425	0.6008772	
770785	0.600311	1,2-alpha-mannosidase IC
2110511	0.5987863	artemin
785795	0.598504	hypothetical protein FLJ12910
120271	0.5982507	hypothetical protein MGC4692
248649	0.5975539	hypothetical protein FLJ13910
2244196	0.5972111	B-cell receptor-associated protein BAP29
66406	0.5966701	hypothetical protein DKFZp762E1312
1537001	0.59528	ESTs
2028949	0.5946445	
40173		KIAA0807 protein
		low molecular mass ubiquinone-binding protein (9.5kD)
100110	0.0002020	10W molocular mass abiquinone binding protein (c.o.c.)
503889	0.5930018	hypothetical protein FLJ10656
781019	0.5918462	paraoxonase 2
196189		cytochrome b-5
1734309	0.5894492	sperm associated antigen 4
		insulin-degrading enzyme
		ATPase inhibitor precursor
		pM5 protein
		H2B histone family, member Q
		ribonucleotide reductase M2 polypeptide
		hexokinase 1
	0.5869599	hippocalcin-like 1
		KIAA0285 gene product
625923		phosphoenolpyruvate carboxykinase 2 (mitochondrial)
2322367	0.5832711	reticulon 4
624667	0.5828923	CGI-92 protein
826363	0.5823873	lysophospholipase II
859228	0.5817344	isocitrate dehydrogenase 1 (NADP+), soluble
		sialidase 3 (membrane sialidase)
209066	0.5794211	serine/threonine kinase 15

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1844765	0.57783	Homo sapiens mRNA; cDNA DKFZp564O1763 (from clone DKFZp564O1763)
191904	0.5770884	BUB3 (budding uninhibited by benzimidazoles 3, yeast) homolog
951233	0.5767874	proteasome (prosome, macropain) subunit, beta type, 3
809357	0.5767382	Bernardinelli-Seip congenital lipodystrophy 2 (seipin)
825740 51532		DKFZp434J1813 protein ADP-ribosylation factor-like 6 interacting protein
203003	0.5743416	non-metastatic cells 4, protein expressed in
2015517 205049		hypothetical protein FLJ22237 protein kinase H11; small stress protein-like protein HSP22
752643 753400 564847 785766 825585 37554 144880	0.5732322 0.573202 0.5731938 0.5722829 0.5706008	group XII secreted phospholipase A2 CGI-204 protein ESTs hypothetical protein tubulin-specific chaperone e hypothetical protein FLJ22353 hypothetical protein from EUROIMAGE 1759349
49273	0.5691181	solute carrier family 27 (fatty acid transporter), member 4
743589	0.568516	ESTs, Weakly similar to T2D3_HUMAN TRANSCRIPTION INITIATION FACTOR TFIID 135 KDA SUBUNIT [H.sapiens]
1616253 1581941 431505 2019223	0.567837 0.5676045 0.5670119 0.5663073	breast carcinoma amplified sequence 1 hypothetical protein FLJ14540 ESTs, Highly similar to A31026 probable membrane receptor protein [H.sapiens] mitochondrial ribosomal protein L17
	0.5648009	
811024		bone marrow stromal cell antigen 2
564492	0.5626905	mitochondrial carrier homolog 2
725841	0.561197	KIAA0662 gene product
701751	0.561139	cut (Drosophila)-like 1 (CCAAT displacement protein)
767798	0.560181	ATX1 (antioxidant protein 1, yeast) homolog 1

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109863 1536006 1845169 343607 212542	0.5590182 0.5588842 0.5588222	epithelial membrane protein 2 ESTs RAB35, member RAS oncogene family AD-015 protein Homo sapiens cDNA FLJ12900 fis, clone NT2RP2004321
2011515 504308 772925 469383 810402	0.5566484 0.5564295 0.5562164 0.5558732	NPD007 protein DKFZP586B0923 protein hypothetical protein FLJ10540 HSPCO34 protein chromosome 8 open reading frame 1 hypothetical protein troponin T1, skeletal, slow
770355	0.5518948	lanosterol synthase (2,3-oxidosqualene-lanosterol cyclase)
813410	0.5513568	polymerase (RNA) II (DNA directed) polypeptide K (7.0kD)
80764 1631735	0.5510626 0.5502891	hypothetical protein Homo sapiens, clone IMAGE:3604336, mRNA, partial cds
625234	0.550211	KDEL (Lys-Asp-Glu-Leu) endoplasmic reticulum protein retention receptor 3
279970	0.5502099	
1518402	0.5497179	KIAA1361 protein
753378	0.548977	hypothetical protein FLJ22649 similar to signal peptidase SPC22/23
594500	0.5474088	
		mitogen-activated protein kinase 13
2055807		phosphoprotein C8FW
773188	0.5454398	nuclear receptor subfamily 1, group D, member 2
126851	0.5454014	hypothetical protein FLJ11160
244764	·	B7 homolog 3
813419	0.5447006	hydroxyacyl-Coenzyme A dehydrogenase, type ll
233349		hypothetical protein FLJ10761
785701		RAB31, member RAS oncogene family
268946	0.5440003	WD40 protein Ciao1
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611443	0.5432426	myoglobin
		hypothetical protein FLJ22087
491465		hypothetical protein FLJ10035
1393018	0.5404243	general transcription factor IIIC, polypeptide 1 (alpha subunit, 220kD)
785840	0.5401209	SEC24 (S. cerevisiae) related gene family, member D
1460110	0.5392177	proteasome (prosome, macropain) subunit, beta type, 5
731308	0.5371204	citrate synthase
301122		extracellular matrix protein 1
66535	0.5363248	ornithine decarboxylase antizyme 2
265103	0.5361177	Homo sapiens mRNA; cDNA DKFZp547M123 (from clone DKFZp547M123)
788334	0.5347204	mitochondrial ribosomal protein L23
813631	0.534683	seven transmembrane protein TM7SF3
1553306	0.5344506	proteasome (prosome, macropain) 26S subunit, non-ATPase,
1461477	0.5343834	Homo sapiens mRNA; cDNA DKFZp586l0324 (from clone DKFZp586l0324)
199645	0.5334951	nicastrin
		chaperonin containing TCP1, subunit 5 (epsilon)
1518890	0.5314577	metallothionein-like 5, testis-specific (tesmin)
756442	0.5313744	P450 (cytochrome) oxidoreductase
		MLN51 protein
		prolactin receptor
		ubiquitin-conjugating enzyme E2 variant 1
		huntingtin (Huntington disease)
795185	0.5276861	xenotropic and polytropic retrovirus receptor
629944	0.5273447	myosin VB
784105	0.5259815	ESTs
1696757		hypothetical protein KIAA1165
139835		UDP-glucose dehydrogenase
1738208	0.5240868	a disintegrin-like and metalloprotease (reprolysin type) with thrombospondin type 1 motif, 4

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593431	0.5235073	ESTs, Moderately similar to CEGT_HUMAN CERAMIDE GLUCOSYLTRANSFERASE [H.sapiens]
825470	0.5218731	frizzled (Drosophila) homolog 5 topoisomerase (DNA) II alpha (170kD) ubiquitin-conjugating enzyme E2N (homologous to yeast UBC13)
376516 774446	0.5195127 0.5181672	interleukin 2 receptor, alpha cell division cycle 4-like
		programmed cell death 9
882484	0.5165756	chaperonin containing TCP1, subunit 7 (eta)
51083	0.5162686	catenin (cadherin-associated protein), delta 2 (neural plakophilin-related arm-repeat protein)
221295	0.5157877	inhibitor of DNA binding 2, dominant negative helix-loop-helix
725978	0.5147267	protein ESTs, Moderately similar to ALU7_HUMAN ALU SUBFAMILY SQ SEQUENCE CONTAMINATION WARNING ENTRY [H.sapiens]
796757	0.5122331	adaptor-related protein complex 3, sigma 1 subunit
150314 302031 293569 838366	_	Ste20-related serine/threonine kinase chromosome 1 open reading frame 21
122147 120749	0.5067128 0.5060444	ESTs, Moderately similar to KIAA1215 protein [H.sapiens]
855563	0.5059683	
487733	0.5057167	NADH dehydrogenase (ubiquinone) 1 beta subcomplex, 2 (8kD, AGGG)
625693	0.5055986	
sd-71385		
	2309073 825470 839682 1517171 376516 774446 1476053 489594 1573946 503851 882484 51083 221295 725978 796757 150314 302031 293569 838366 122147 120749 855563 487733 625693	2309073

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813751	0.5051993	sialyltransferase 4C (beta-galactosidase alpha-2,3-
010701		sialytransferase)
842825	0.5047729	G1 to S phase transition 1
323693		adaptor-related protein complex 1, sigma 1 subunit
202514	0.5045411	DNA (cytosine-5-)-methyltransferase 3 alpha
1517749	0.5045134	ESTs
124331		cleavage and polyadenylation specific factor 5, 25 kD subunit
1474424	0.5021308	Homo sapiens cDNA FLJ12758 fis, clone NT2RP2001328
1848977	0.5015371	glycerol kinase
454896		DnaJ (Hsp40) homolog, subfamily A, member 2
1702742	0.5011725	solute carrier family 7 (cationic amino acid transporter, y+ system), member 5
746163	0.5009715	ESTs, Weakly similar to ALU1_HUMAN ALU SUBFAMILY J SEQUENCE CONTAMINATION WARNING ENTRY
		[H.sapiens]
810734	0.4998276	polymerase (DNA-directed), delta 4
725284	0.4992842	phosphorylase kinase, gamma 2 (testis)
52103	0.4988158	hypothetical protein FLJ23045
739109	0.4986821	adaptor-related protein complex 2, sigma 1 subunit
488505	0.4981852	accessory proteins BAP31/BAP29
365060	0.4976091	RAB11A, member RAS oncogene family
41569	0.4974126	hypothetical protein FLJ12650
81336		uteroglobin
700792	0.4949149	cyclin-dependent kinase inhibitor 3 (CDK2-associated dual specificity phosphatase)
1592715	0.4944665	Homer, neuronal immediate early gene, 3
142586	0.4942961	MCT-1 protein
261472	0.4941066	putative nuclear protein ORF1-FL49
246800	0.4927231	hypothetical protein FLJ10803
124298	0.4922937	microsomal glutathione S-transferase 3
1898619	0.4916939	hypothetical protein MGC15737
739126	0.4905859	tissue specific transplantation antigen P35B
376875	0.490328	flavin containing monooxygenase 1
358162	0.4901786	protein predicted by clone 23627
590759	0.4895021	sterol-C4-methyl oxidase-like
sd-71385		

1947647 120271 488642	0.4886584	CGI-147 protein hypothetical protein MGC4692 ESTs, Weakly similar to I38022 hypothetical protein [H.sapiens]
1492468	0.4878466	DEME-6 protein
345538	0.4875796	cathepsin L
344091	0.4871044	ESTs
123441	0.4862068	ribosomal protein L7a
202901	0.48608	vav 2 oncogene
757489	0.4859039	tubulin, alpha 2
49117		KIAA0215 gene product
812994		retinoid X receptor, alpha
731023		WD repeat domain 5
814899	0.4840969	BCL2/adenovirus E1B 19kD-interacting protein 3-like
756666	0.4835569	protein phosphatase 1, catalytic subunit, alpha isoform
2018337	0.4833207	glucosidase, beta; acid (includes glucosylceramidase)
1568825	0.4829928	Arg/Abl-interacting protein ArgBP2
869375	0.4827093	isocitrate dehydrogenase 2 (NADP+), mitochondrial
595213	0.4824381	hypothetical protein
	0.4824256	putative ankyrin-repeat containing protein
73009	0.4822587	ESTs, Weakly similar to A43932 mucin 2 precursor, intestinal [H.sapiens]
70606	0.481356	• • •
358456	0.4812268	Sec61 gamma
810762		SNARE protein
823930	0.4800704	actin related protein 2/3 complex, subunit 1A (41 kD)
2020898	0.4797032	procollagen-lysine, 2-oxoglutarate 5-dioxygenase 3
564981	0.4782207	ESTs
1637282	0.4777562	hexokinase 2
855749	0.476968	triosephosphate isomerase 1
744944	0.4769116	myosin VI
263727	0.4767486	DNA segment, single copy probe LNS-CAI/LNS-CAII (deleted in polyposis
470092	0.476018	like-glycosyltransferase
782513		interferon, alpha-inducible protein (clone IFI-6-16)

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897813	0.4749348	polyadenylate binding protein-interacting protein 1	
141852	0.4748862	purinergic receptor P2Y, G-protein coupled, 2	
855800 41356	0.4748367 0.4746307	prolyl endopeptidase protein phosphatase 2, regulatory subunit B (B56), alpha isoform	
		WD repeat domain 15	
813387		KIAA0143 protein diaphorase (NADH/NADPH) (cytochrome b-5 reductase)	
358267	0.4719017	EST, Moderately similar to AF119917 63 PRO2831 [H.sapiens]	
2018527	0.4717524	dolichyl-phosphate mannosyltransferase polypeptide 3	
785707 140635 814306	0.4708376 0.4706227 0.4705205 0.4702303	hypothetical protein MGC11115 protein regulator of cytokinesis 1 ESTs tumor protein D52 RAB31, member RAS oncogene family Homo sapiens, Similar to RIKEN cDNA 5830420C20 gene, clone IMAGE:3633379, mRNA, partial cds	
377384	0.4698441	nuclear receptor subfamily 2, group F, member 2	
183200	0.4697062	fumarylacetoacetate hydrolase (fumarylacetoacetase)	
345423 204686		DKFZP564M112 protein FXYD domain-containing ion transport regulator 1 (phospholemman)	
428582 210862 51741	0.4683996 0.4678807	hypothetical protein FLJ20296 acyl-Coenzyme A oxidase 1, palmitoyl GTP-binding protein	
840865	0.4670993	deiodinase, iodothyronine, type I myristoylated alanine-rich protein kinase C substrate (MARCKS, 80K-L)	
278531 60565 1639531 346942 884498	0.4663008 0.4659865 0.4658349 0.4657601 0.4648731	cytochrome c oxidase subunit VIc lethal giant larvae (Drosophila) homolog 2	

345787	0.4647372	highly expressed in cancer, rich in leucine heptad repeats
825296	0.4646099	low density lipoprotein receptor defect C complementing
2248488	0.4645687	ems1 sequence (mammary tumor and squamous cell carcinoma-associated (p80/85 src substrate)
824962	0.4638758	karyopherin alpha 2 (RAG cohort 1, importin alpha 1)
843054	0.4637278	KIAA1533 protein
140289	0.4624364	hypothetical protein
1558233	0.4622449	ESTs
265592	0.4615543	activated RNA polymerase II transcription cofactor 4
767761	0.4608709	DKFZP434B168 protein
770766	0.4603527	DKFZP564C1940 protein
814209	0.4602081	ESTs
773674	0.4601667	oncogene TC21
703707	0.4599434	aspartate beta-hydroxylase
951216	0.4594172	NADH dehydrogenase (ubiquinone) 1 beta subcomplex, 10 (22kD, PDSW)
139354	0.4582974	hypothetical protein
79710		KIAA0174 gene product
823574		endosulfine alpha
845363	0.4570873	non-metastatic cells 1, protein (NM23A) expressed in
283751	0.4567022	cortistatin
810156	0.4561539	deoxythymidylate kinase (thymidylate kinase)
1416782	0.4558415	creatine kinase, brain
810609	0.4551167	hypothetical protein PP1226
770675	0.454729	Homo sapiens cDNA: FLJ21323 fis, clone COL02374
626318	0.4529715	ubinuclein 1
38356		cytochrome P450, subfamily 46 (cholesterol 24-hydroxylase)
705064	0.4497555	transforming, acidic coiled-coil containing protein 3
127646	0.4493747	
		hypothetical protein DC42
		cleavage stimulation factor, 3' pre-RNA, subunit 3, 77kD
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489823	0.4468978	COX17 (yeast) homolog, cytochrome c oxidase assembly protein
469151	0.4462659	eukaryotic translation initiation factor 2, subunit 2 (beta, 38kD)
41826 47853	0.4456463 0.445149	ESTs aldehyde dehydrogenase 4 family, member A1
144740	0.4449229	similar to phosphatidylcholine transfer protein 2
726658	0.4446657	non-metastatic cells 3, protein expressed in
325515 815781 746190 2016194 1584287 147834 154610 754653	0.4441245 0.4430243 0.4417536 0.4413331 0.4412795 0.4410751 0.4405873	cadherin, EGF LAG seven-pass G-type receptor 3, flamingo (Drosophila) homolog
742581	0.440367	Homo sapiens cDNA FLJ10366 fis, clone NT2RM2001420
2306987	0.4402912	secreted and transmembrane 1
878406	0.4396935	
430614		2,3-bisphosphoglycerate mutase
		cell division cycle 25C
35821 <i>7</i> 1636669	0.4385343 0.4384707	
83358	0.4380365	
1325816	0.4379227	polymerase (RNA) II (DNA directed) polypeptide L (7.6kD)
824524	0 4375695	UDP-galactose transporter related
795805 743977 785793	0.4370764 0.4368155	KIAA0332 protein Homo sapiens mRNA for TL132 capping protein (actin filament) muscle Z-line, alpha 1

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782503 753897 767289	0.4339171 0.4334388 0.4324066 0.4323485 0.4323117 0.4321289	hypothetical protein FLJ22056 thioredoxin reductase 1 fatty acid desaturase 1 autocrine motility factor receptor hypothetical protein FLJ10055 interferon regulatory factor 7 Homo sapiens, clone MGC:19613 IMAGE:3833049, mRNA, complete cds ESTs, Weakly similar to T2D3_HUMAN TRANSCRIPTION INITIATION FACTOR TFIID 135 KDA SUBUNIT [H.sapiens]
41698 1635665	0.4310061 0.430087	progesterone binding protein Homo sapiens, RIKEN cDNA 2010100O12 gene, clone MGC:14813 IMAGE:4133274, mRNA, complete cds
	0.4298376 0.4289711 0.4280476	hypothetical protein MGC2771 BLu protein Homo sapiens chromosome 19, BAC CIT-HSPC_204F22 (BC228680), complete sequence; contains bacterial insertion element
	0.4277815 0.4277651	ESTs, Weakly similar to T26581 hypothetical protein Y32B12A.3 - Caenorhabditis elegans [C.elegans]
731080 649084 795498 897770 2250839	0.4275738 0.4270889 0.4270685	hypothetical protein FLJ12661 carbonic anhydrase XI putative transmembrane protein androgen receptor (dihydrotestosterone receptor; testicular feminization; spinal and bulbar muscular atrophy; Kennedy
503866	0.4255725 0.424655 0.4242762 0.4242722 0.4230404	c-myc binding protein hypothetical protein N-myc downstream regulated pyruvate dehydrogenase (lipoamide) beta hypothetical protein MGC861 sperm autoantigenic protein 17 Homo sapiens, Similar to CG7083 gene product, clone MGC:10534 IMAGE:3957147, mRNA, complete cds
292936 sd-71385	0.4217115	hypothetical protein FLJ10468

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823940 414992		transducer of ERBB2, 1 K562 cell-derived leucine-zipper-like protein 1
346134	0.4209894	calcium-regulated heat-stable protein (24kD)
	0.4177687 0.4171445 0.4167519	rabaptin-5 oncostatin M receptor synaptotagmin-like 2
1762111	0.4160469	natriuretic peptide receptor C/guanylate cyclase C (atrionatriuretic peptide receptor C)
509588	0.4157059	TATA box binding protein (TBP)-associated factor, RNA polymerase II, J, 20kD
327506	0.4152106	Homo sapiens mRNA full length insert cDNA clone EUROIMAGE 327506
744047	0.4151137	polo (Drosophia)-like kinase
221499		KIAA0508 protein
745394	0.4147627	Homo sapiens cDNA: FLJ23249 fis, clone COL04196
149539	0.4138942	KIAA1700
823907	0.4135955	hypothetical protein FLJ10511
66317	0.4132405	H1 histone family, member 2
295986	0.413087	emopamil-binding protein (sterol isomerase)
565235	0.4124667	spermine synthase
825386	0.4121622	ATP synthase, H+ transporting, mitochondrial F1F0, subunit d
250313	0.4121296	ESTs
72050	0.4114235	chloride channel, nucleotide-sensitive, 1A
417801	0.4112944	mitochondrial ribosomal protein L27
841501	0.4111707	KIAA0102 gene product
813707	0.410787	regulator of G-protein signalling 16
745083	0.4107735	ubiquitin specific protease 18
244974	0.4105439	hypothetical protein FLJ22875
26883	0.4103824	protein kinase (cAMP-dependent, catalytic) inhibitor beta

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280249 74738		Kruppel-like factor 7 (ubiquitous) Homo sapiens, clone IMAGE:3535294, mRNA, partial cds
1753497 204299 358609	0.4073087	ovo (Drosophila) homolog-like 1 replication protein A3 (14kD) NADH dehydrogenase (ubiquinone) flavoprotein 3 (10kD)
290101 2306221	0.4071477 0.4068933	ESTs wingless-type MMTV integration site family, member 10B
726439 431805 111362	0.405447	CGI-143 protein granulin
472103		soc-2 (suppressor of clear, C.elegans) homolog
788444 1632252	0.4042598 0.404193	KIAA0033 protein complement component 1, q subcomponent, alpha polypeptide
150118 594226		hypothetical protein DKFZp434F054 Homo sapiens cDNA FLJ14459 fis, clone HEMBB1002409
1499828	0.4030396	fucosyltransferase 1 (galactoside 2-alpha-L-fucosyltransferase, Bombay phenotype included)
810939 841621	0.40256 0.4020253	hypothetical protein FLJ22169 ESTs, Weakly similar to JE0350 Anterior gradient-2 [H.sapiens]
200144 486110 292213	0.4019827 0.4017126 0.4014458	
1631132 669379	0.401216 0.4011648	peptide transporter 3 Homo sapiens BAC clone RP11-505D17 from 7p22-p21
773685	0.4009493	phosphodiesterase 4D interacting protein (myomegalin)
810411 1583198	0.400096 0.39981	hypothetical protein FLJ13222 ESTs, Weakly similar to S65824 reverse transcriptase homolog [H.sapiens]
84295 757328 35626	0.3989864 0.3985331 0.39822	interleukin 1 receptor antagonist hypothetical protein FLJ22678 Homo sapiens cDNA FLJ14201 fis, clone NT2RP3002955

1587863	0.3980812	acetyl-Coenzyme A acyltransferase 1 (peroxisomal 3-oxoacyl-Coenzyme A thiolase)
713862		ubiquitin-protein isopeptide ligase (E3)
343731 43977	0.3962504 0.3961848	KIAA0182 protein
78869	0.3954821	cell membrane glycoprotein, 110000M(r) (surface antigen)
756595	0.395121	S100 calcium-binding protein A10 (annexin II ligand, calpactin I, light polypeptide (p11))
701115	0.3936994	
81599	0.3932769	ubiquitin specific protease 14 (tRNA-guanine transglycosylase)
1636844		ring finger protein 14
703739	0.392298	nuclear cap binding protein subunit 1, 80kD
279633	0.3907503	ESTs, Highly similar to I38759 zinc finger/leucine zipper protein [H.sapiens]
122091	0.3903482	, t
786067 1909574	0.390131	cell division cycle 25B mitochondrial ribosomal protein S11
263013	0.3894536	•
		hydroxylase) 2
491001	0.3884144	
455275 85450	0.3868399	hypothetical protein FLJ23469 acyl-Coenzyme A oxidase 2, branched chain
773286	0.3001343	solute carrier family 9 (sodium/hydrogen exchanger), isoform 3 regulatory factor 1
366156	0.3861394	Homo sapiens cDNA FLJ14028 fis, clone HEMBA1003838
49630	0.3858335	calcium channel, voltage-dependent, L type, alpha 1D subunit
121251	0.385792	hypothetical protein MGC5576
2322223	0.3853156	small nuclear ribonucleoprotein polypeptide A
276915	0.3851081	DNA (cytosine-5-)-methyltransferase 3 beta
1591264	0.3849595	transaldolase 1
		HIV TAT specific factor 1
489657	0.3847527	tryptophan rich basic protein

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810947	0.3833137	LIS1-interacting protein NUDE1, rat homolog
298417 154707		trefoil factor 3 (intestinal) MpV17 transgene, murine homolog, glomerulosclerosis
		CGI-82 protein BCL2-interacting killer (apoptosis-inducing)
1455394 769942 429387 768377 214996 1557637 1517595	0.3810371 0.3810004 0.3809674 0.380548 0.3805285 0.3804132 0.3801713 0.3785047	cytochrome c kinesin-like 4 chimerin (chimaerin) 2 activity-dependent neuroprotective protein ESTs
76196 2017721	0.3781067	[H.sapiens]
		homeo box D8 FGFR1 oncogene partner
1649374	0.3771383	homogentisate 1,2-dioxygenase (homogentisate oxidase)
143426 1412245 786265 266500	0.3764754 0.3764075 0.3762847 0.3760397 0.3758559 0.3749416 0.3744208 0.3741516	Human clone 137308 mRNA, partial cds hypothetical protein R33729_1 ras homolog gene family, member B carboxypeptidase A2 (pancreatic) KIAA0750 gene product ESTs protein kinase C, delta DKFZP586A0522 protein Homo sapiens, Similar to RIKEN cDNA 2210021G21 gene, clone MGC:14859 IMAGE:3621871, mRNA, complete cds
454339	0.3738929	thiopurine S-methyltransferase

810316	0.3732635	very long-chain acyl-CoA synthetase; lipidosin
772898	0.3727398 0.3725734 0.372064	ALL1-fused gene from chromosome 1q KIAA0391 gene product hypothetical protein FLJ21213 ribosomal protein S15a Homo sapiens cDNA FLJ14028 fis, clone HEMBA1003838
289505	0.3714361	ESTs, Moderately similar to S65657 alpha-1C-adrenergic receptor splice form 2 [H.sapiens]
	0.3708058	phosphoglycerate mutase 1 (brain) ESTs Homo sapiens cDNA FLJ13975 fis, clone Y79AA1001585
1679942 566443		KIAA1053 protein Homo sapiens cDNA FLJ12793 fis, clone NT2RP2002033
302996 726699		chloride intracellular channel 3 ESTs, Weakly similar to AAB47496 NG5 [H.sapiens]
2094232 2549634 177827 852829 207794 823909	0.3692338 0.3691459 0.3691079 0.3679245 0.3675927	chromosome 1 open reading frame 12 activator of S phase kinase synaptotagmin VII karyopherin alpha 3 (importin alpha 4) nuclear factor (erythroid-derived 2), 45kD
743810 810039 53039	0.3662086	hypothetical protein MGC2577 defender against cell death 1 carbohydrate (keratan sulfate Gal-6) sulfotransferase 1
809727 26171 1570663 179212	0.3656056	unc-51 (C. elegans)-like kinase 1 KIAA0856 protein FK506-binding protein 4 (59kD) ESTs, Moderately similar to T12539 hypothetical protein DKFZp434J154.1 [H.sapiens]
292996	0.36439	tyrosine 3-monooxygenase/tryptophan 5-monooxygenase activation protein, eta polypeptide
259950 815794 965223 sd-71385		hypothetical protein FLJ14991 nucleobindin 2 thymidine kinase 1, soluble

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592801	0.3638731	serine palmitoyltransferase, long chain base subunit 2
42831 788641		N-terminal kinase-like adaptor-related protein complex 1, sigma 2 subunit
813845 52076 154493 1519013	0.3633002 0.3629466	RNA, U transporter 1 neuroblastoma (nerve tissue) protein interferon-induced protein 75, 52kD Homo sapiens, clone IMAGE:3537447, mRNA, partial cds
1434905 825282 322617	0.3624112	homeo box B7 DKFZP586L0724 protein v-ral simian leukemia viral oncogene homolog B (ras related; GTP binding protein)
814353	0.3622592	phorbol-12-myristate-13-acetate-induced protein 1
2017917	0.360911	Homo sapiens cDNA: FLJ23371 fis, clone HEP16068, highly similar to HSTFIISH Homo sapiens mRNA for transcription elongation factor TFIIS
813616 1846982 505289		FK506-binding protein like inhibin, beta C angiotensin II, type I receptor-associated protein
1565079 795401	0.358728 0.3573481	B-cell linker diacylglycerol O-acyltransferase (mouse) homolog
781047	0.357207	budding uninhibited by benzimidazoles 1 (yeast homolog)
346696 809466	0.3570344 0.3568464	TEA domain family member 4 DNA segment on chromosome 19 (unique) 1177 expressed sequence
743220 131268 119133	0.3566593 0.3564734 0.3564449	hypothetical protein FLJ12517 growth factor receptor-bound protein 14
242706	0.3562469	Homo sapiens, Similar to RIKEN cDNA 5730494N06 gene, clone MGC:13349 IMAGE:4249231, mRNA, complete cds
66599	0.3561699	N-acetyltransferase 1 (arylamine N-acetyltransferase)
46716 sd-71385	0.3557556	ESTs

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271899	0.355689	Ser-Thr protein kinase related to the myotonic dystrophy protein kinase
25440	0.3554359	
1899312 783681	0.3551595 0.354881	ESTs upstream regulatory element binding protein 1
131091 300099 588436 2069602 810959		butyrate-induced transcript 1
649977		Homo sapiens clone CDABP0014 mRNA sequence
345069 767487	0.3532976	nuclear factor (erythroid-derived 2)-like 3 ariadne (Drosophila) homolog, ubiquitin-conjugating enzyme E2-binding protein, 1
824943 744994 773381		hypothetical protein hypothetical protein FLJ12242
1568561 2028916		BCL2-like 1 Homo sapiens mRNA for Hmob33 protein, 3' untranslated region
753457	0.3515361	NADH dehydrogenase (ubiquinone) Fe-S protein 1 (75kD) (NADH-coenzyme Q reductase)
2161427 1947381 82421 504461 754625 795543	0.3509235 0.3506778 0.3506753 0.3503938	progesterone receptor hypothetical protein FLJ22329 zinc finger protein 6 (CMPX1) opsin 3 (encephalopsin) ATPase, Class II, type 9A thioredoxin peroxidase (antioxidant enzyme)
172785	0.3493683	KIAA0535 gene product NAG-5 protein ESTs, Weakly similar to YK54_YEAST HYPOTHETICAL 18.4 KD PROTEIN IN SIS2-MTD1 INTERGENIC REGION [S.cerevisiae]
321354 sd-71385	0.3489273	mitochondrial ribosomal protein L15

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878815 741769 487831	0.3475255	ADP-ribosylation factor 3 polymerase (DNA directed), beta Homo sapiens cDNA FLJ14059 fis, clone HEMBB1000573
754046	0.3464097	DNA segment on chromosome X (unique) 9879 expressed sequence
344988 530093 823615	0.3464047 0.3463461 0.3462131	myelin protein zero-like 1 Homo sapiens cDNA: FLJ21245 fis, clone COL01184
144881 128695	0.3461767 0.3455381	calumenin ESTs, Weakly similar to I38344 titin, cardiac muscle [H.sapiens]
884511 128711		cytochrome c oxidase subunit VIIb anillin (Drosophila Scraps homolog), actin binding protein
470035 1568126	0.3436575 0.3434836	prokineticin 1 precursor solute carrier family 37 (glycerol-3-phosphate transporter), member 1
588911	0.3427563	2',5'-oligoadenylate synthetase 1 (40-46 kD)
35191 814989	+ - + - · + - ·	stromal cell-derived factor 2 protein phosphatase 1G (formerly 2C), magnesium-dependent, gamma isoform
73638	0.3425644	protein tyrosine phosphatase type IVA, member 2
1605178 214068 755599	0.3414185	amino acid transporter system A1 GATA-binding protein 3 interferon induced transmembrane protein 1 (9-27)
782439	0.3413092	ATP synthase, H+ transporting, mitochondrial F0 complex,
595070	0.3408109	subunit e stress-associated endoplasmic reticulum protein 1; ribosome associated membrane protein 4
183440 774502	0.3406314 0.3402891	arylsulfatase A protein tyrosine phosphatase, non-receptor type 12
235986	0.338857	wingless-type MMTV integration site family, member 11
289936	0.3382708	solute carrier family 7, (cationic amino acid transporter, y+ system) member 11
824108 sd-71385	0.3381616	SCAN domain-containing 1

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26617	0.3371923	activated leucocyte cell adhesion molecule
810497	0.3367408	ESTs, Weakly similar to A35363 synapsin I splice form a [H.sapiens]
40042	0.3362557	hypothetical protein FLJ10747
811142		phosphoinositide-3-kinase, regulatory subunit, polypeptide 2 (p85 beta)
2046679	0.3358043	ËSTs
487444	0.3357395	cyclic AMP phosphoprotein, 19 kD
78736	0.3354693	Homo sapiens clone 24877 mRNA sequence
1637302	0.3353374	DNAJ domain-containing
825606	0.335057	kinesin-like 1
76362	0.3349202	spectrin, alpha, non-erythrocytic 1 (alpha-fodrin)
1883028	0.3345488	Homo sapiens mRNA; cDNA DKFZp434J1912 (from clone DKFZp434J1912)
789383	0.3341667	cAMP responsive element modulator
302549		basic transcription element binding protein 1
002010	0.00.000	paste transcription cromonium annam g process.
309288	0.334016	replication factor C (activator 1) 4 (37kD)
712314	0.3338146	myosin regulatory light chain interacting protein
241677	0.3332521	Homo sapiens, clone MGC:18110 IMAGE:4152745, mRNA,
000050	0.0004004	complete cds
898253	0.3331924	reticulocalbin 2, EF-hand calcium binding domain
770992	0.3331315	
81203	0.3328573	paraoxonase 3
416436	0.3325571	mitochondrial ribosomal protein L24
1660666	0.332029	carbonic anhydrase VB, mitochondrial
811582	0.3319191	golgi phosphoprotein 2
2054122	0.3318413	solute carrier family 11 (proton-coupled divalent metal ion
		transporters), member 3
837864	0.3314764	progestin induced protein
1632247	0.3312131	hypothetical protein FLJ23436
298143	0.3307258	
		hypothetical protein FLJ20297
1600281	0.3304323	nucleolar protein 3 (apoptosis repressor with CARD domain)
814270	0.3300091	polymyositis/scleroderma autoantigen 1 (75kD)

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415589	0.3296266	Homo sapiens clone PP1498 unknown mRNA
1912951 726637		uterine-derived 14 kDa protein t-complex-associated-testis-expressed 1-like
1843843		ephrin-A1 KIAA1304 protein ATPase, Cu++ transporting, beta polypeptide (Wilson disease)
135640 279720		syntaxin 3A ESTs, Moderately similar to A47582 B-cell growth factor precursor [H.sapiens]
724888	0.3272422	cytochrome P450, subfamily IVB, polypeptide 1
811808	0.326728	nucleoside diphosphate kinase type 6 (inhibitor of p53-induced apoptosis-alpha)
131566 154466		hypothetical protein MGC11061 STIP1 homology and U-Box containing protein 1
1692195 785368	0.3254467 0.3254362	smg GDS-ASSOCIATED PROTEIN PDZ-binding kinase; T-cell originated protein kinase
	0.32462320.32439840.3242547	Homo sapiens, clone MGC:3182 IMAGE:3356293, mRNA, complete cds
		IMAGE:4130879, mRNA, partial cds
855406	0.3241172	Homo sapiens clone TCCCIA00176 mRNA sequence
1500542 269606 74566	0.3228485 0.3228482 0.3226444	CGI-107 protein regulator of G-protein signalling 11 N-methylpurine-DNA glycosylase exportin 1 (CRM1, yeast, homolog) Homo sapiens mRNA; cDNA DKFZp586H0924 (from clone DKFZp586H0924)
180785 824479 842863 1708055 sd-71385	0.3225263 0.3225116	KIAA0726 gene product exonuclease NEF-sp N-myc downstream regulated glioblastoma overexpressed

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796542	-1.2625621	interleukin 11 receptor, alpha ets variant gene 5 (ets-related molecule) Homo sapiens mRNA; cDNA DKFZp564J0323 (from clone DKFZp564J0323)
753071	-1.2529315	Homo sapiens cDNA: FLJ22528 fis, clone HRC12825
		folate receptor 1 (adult) hypothetical protein FLJ21044 similar to Rbig1
		crystallin, alpha B Homo sapiens mRNA; cDNA DKFZp564H1916 (from clone DKFZp564H1916)
490023 298122	-1.2114805 -1.1962478	lipoma HMGIC fusion partner-like 2 hypothetical protein MGC2648 frizzled (Drosophila) homolog 7 caspase 7, apoptosis-related cysteine protease
160192	-1.1751869	ESTs, Weakly similar to 2004399A chromosomal protein
22917	-1.1590598	[H.sapiens] Homo sapiens mRNA; cDNA DKFZp761M0111 (from clone DKFZp761M0111)
		nGAP-like protein Homo sapiens cDNA: FLJ22667 fis, clone HSI08385
289760	-1.1390686	ESTs, Highly similar to T00391 hypothetical protein KIAA0612 [H.sapiens]
1556433 838478	-1.1222978 -1.1175047	hypothetical protein GRO3 oncogene neurocalcin delta myosin regulatory light chain 2, smooth muscle isoform
811088 1554167 67741 2106144 2056139	-1.0851546 -1.0818834 -1.0705249 -1.067765 -1.0673271	serum amyloid A4, constitutive ephrin-B3 hypothetical protein FLJ14529 PP2135 protein regulated in glioma LIM domain protein latent transforming growth factor beta binding protein 2

345034	-1.0600859	small inducible cytokine subfamily B (Cys-X-Cys), member 14 (BRAK)
666879	-1.0463245	
130201	-1.0401997	intercellular adhesion molecule 2
878836	-1.037352	secretory granule, neuroendocrine protein 1 (7B2 protein)
52419	-1 0368509	Friedreich ataxia region gene X123
		runt-related transcription factor 3
		suppression of tumorigenicity 5
132857	-1.0084069	Homo sapiens mRNA; cDNA DKFZp586N1323 (from clone
		DKFZp586N1323)
529843	-1.0036824	ESTs, Moderately similar to JC5238 galactosylceramide-like
		protein, GCP [H.sapiens]
2504881	-1.000863	signal transducer and activator of transcription 5A
69002	-0.9960916	PPAR(gamma) angiopoietin related protein
841308	-0.9953716	myosin, light polypeptide kinase
488404		Homo sapiens clone TUA8 Cri-du-chat region mRNA
277571	-0.9841522	KIAA1706 protein
		Homo sapiens mRNA; cDNA DKFZp434C1714 (from clone
		DKFZp434C1714); partial cds
897963	-0.9775616	phosphatidic acid phosphatase type 2A
		Homo sapiens mRNA; cDNA DKFZp586M2022 (from clone
		DKFZp586M2022)
196435	-0.9665947	ESTs
140574	-0.9652218	small inducible cytokine subfamily D (Cys-X3-Cys), member 1
		(fractalkine, neurotactin)
740620	-0.96496	tropomyosin 2 (beta)
		KIAA0194 protein
1635320	-0.9586226	amiloride-sensitive cation channel 2, neuronal
416959	-0.957248	nuclear factor I/B
814798		aldehyde dehydrogenase 1 family, member A3
000074	0.0400440	CDADC like 1 (most) hovin
		SPARC-like 1 (mast9, hevin)
303109	-0.5414002	purinergic receptor (family A group 5)



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270826	-0.9389651	Homo sapiens cDNA FLJ13329 fis, clone OVARC1001795
66491 814316	-0.9280245	ribosomal protein L13
811149	-0.9216006	chromosome 9 open reading frame 3 beta-site APP-cleaving enzyme
	-0.9190244	• ,
416676 248631		pellino (Drosophila) homolog 1 aminomethyltransferase (glycine cleavage system protein T)
76182 756708		hypothetical protein DKFZp761F241 potassium intermediate/small conductance calcium-activated channel, subfamily N, member 4
712401	-0.9068788	phosphoinositide-3-kinase, catalytic, delta polypeptide
625399	-0.9066615	hypothetical protein similar to beta-transducin family
	-0.893161 -0.8910356	kinesin family member C3 ESTs
		growth arrest-specific 6
133236 415816	-0.8887425	RNA binding motif protein, X chromosome ESTs
323780	-0.8879854	Homo sapiens cDNA FLJ11177 fis, clone PLACE1007402
		hypothetical protein FLJ22297 KIAA0668 protein
		tweety (Drosophila) homolog 1
	-0.8800259	
		osteoblast specific factor 2 (fasciclin I-like) eukaryotic translation elongation factor 1 alpha 1
		forkhead box O1A (rhabdomyosarcoma)
		Down syndrome critical region gene 1 NGFI-A binding protein 1 (EGR1 binding protein 1)
882248 770935	-0.8700731 -0.8700104	transgelin hypothetical protein FLJ13511
41208	-0.865227	bone morphogenetic protein 1
768571	-0.8572018	SRY (sex determining region Y)-box 8
1.54000		

781017	-0.8526926	early growth response 2 (Krox-20 (Drosophila) homolog)
	-0.848137 -0.8444322	ATP synthase, H+ transporting, mitochondrial F1 complex, alpha subunit, isoform 1, cardiac muscle
814826	-0.8443265	ESTs
767164	-0.842198	ESTs, Weakly similar to MUC2_HUMAN MUCIN 2 PRECURSOR [H.sapiens]
71087	-0.8416951	v-maf musculoaponeurotic fibrosarcoma (avian) oncogene family, protein F
283023	-0.8413949	chemokine (C-X3-C) receptor 1
	-0.8405669	
		hypothetical protein MGC3232
142259	-0.8376566	tumor necrosis factor alpha-inducible cellular protein containing leucine zipper domains; Huntingtin interacting protein L; transcrption factor IIIA-interacting protein
990881	-0.8335353	·
970590	-0.8333968	Homo sapiens mRNA; cDNA DKFZp434A115 (from clone DKFZp434A115)
470393	-0.830751	matrix metalloproteinase 7 (matrilysin, uterine)
	-0.830751 -0.8306579	
772913	-0.8306579	
772913 741139	-0.8306579	calreticulin eyes absent (Drosophila) homolog 2
772913 741139 27769	-0.8306579 -0.8299902	calreticulin eyes absent (Drosophila) homolog 2 ESTs
772913 741139 27769 544639 1587710	-0.8306579 -0.8299902 -0.8298821 -0.8275232 -0.8252065	calreticulin eyes absent (Drosophila) homolog 2 ESTs ESTs period (Drosophila) homolog 1
772913 741139 27769 544639 1587710 757191	-0.8306579 -0.8299902 -0.8298821 -0.8275232 -0.8252065 -0.8247438	calreticulin eyes absent (Drosophila) homolog 2 ESTs ESTs period (Drosophila) homolog 1 ESTs
772913 741139 27769 544639 1587710 757191 80344	-0.8306579 -0.8299902 -0.8298821 -0.8275232 -0.8252065 -0.8247438 -0.8243418	calreticulin eyes absent (Drosophila) homolog 2 ESTs ESTs period (Drosophila) homolog 1 ESTs interleukin 7 receptor
772913 741139 27769 544639 1587710 757191 80344 1635062	-0.8306579 -0.8299902 -0.8298821 -0.8275232 -0.8252065 -0.8247438 -0.8243418 -0.8243024	calreticulin eyes absent (Drosophila) homolog 2 ESTs ESTs period (Drosophila) homolog 1 ESTs interleukin 7 receptor DKFZP586A011 protein
772913 741139 27769 544639 1587710 757191 80344 1635062	-0.8306579 -0.8299902 -0.8298821 -0.8275232 -0.8252065 -0.8247438 -0.8243418 -0.8243024	calreticulin eyes absent (Drosophila) homolog 2 ESTs ESTs period (Drosophila) homolog 1 ESTs interleukin 7 receptor
772913 741139 27769 544639 1587710 757191 80344 1635062 119290	-0.8306579 -0.8299902 -0.8298821 -0.8275232 -0.8252065 -0.8247438 -0.8243418 -0.8243024 -0.8224756	calreticulin eyes absent (Drosophila) homolog 2 ESTs ESTs period (Drosophila) homolog 1 ESTs interleukin 7 receptor DKFZP586A011 protein cortic al thymocyte receptor (X. laevis CTX) like
772913 741139 27769 544639 1587710 757191 80344 1635062 119290	-0.8306579 -0.8299902 -0.8298821 -0.8275232 -0.8252065 -0.8247438 -0.8243418 -0.8243024 -0.8224756	calreticulin eyes absent (Drosophila) homolog 2 ESTs ESTs period (Drosophila) homolog 1 ESTs interleukin 7 receptor DKFZP586A011 protein cortic al thymocyte receptor (X. laevis CTX) like serum amyloid A1
772913 741139 27769 544639 1587710 757191 80344 1635062 119290 161456 1909935	-0.8306579 -0.8299902 -0.8298821 -0.8275232 -0.8252065 -0.8247438 -0.8243418 -0.8243024 -0.8224756 -0.8215502	calreticulin eyes absent (Drosophila) homolog 2 ESTs ESTs period (Drosophila) homolog 1 ESTs interleukin 7 receptor DKFZP586A011 protein cortic al thymocyte receptor (X. laevis CTX) like serum amyloid A1 ESTs
772913 741139 27769 544639 1587710 757191 80344 1635062 119290 161456 1909935	-0.8306579 -0.8299902 -0.8298821 -0.8275232 -0.8252065 -0.8247438 -0.8243418 -0.8243024 -0.8224756 -0.8215502 -0.8169271 -0.8163443	calreticulin eyes absent (Drosophila) homolog 2 ESTs ESTs period (Drosophila) homolog 1 ESTs interleukin 7 receptor DKFZP586A011 protein cortic al thymocyte receptor (X. laevis CTX) like serum amyloid A1 ESTs
772913 741139 27769 544639 1587710 757191 80344 1635062 119290 161456 1909935 1257131 868396	-0.8306579 -0.8299902 -0.8298821 -0.8275232 -0.8252065 -0.8247438 -0.8243418 -0.8243024 -0.8224756 -0.8215502 -0.8169271 -0.8163443 -0.808677	calreticulin eyes absent (Drosophila) homolog 2 ESTs ESTs period (Drosophila) homolog 1 ESTs interleukin 7 receptor DKFZP586A011 protein cortic al thymocyte receptor (X. laevis CTX) like serum amyloid A1 ESTs ESTs ESTs tyrosine 3-monooxygenase/tryptophan 5-monooxygenase
772913 741139 27769 544639 1587710 757191 80344 1635062 119290 161456 1909935 1257131 868396	-0.8306579 -0.8299902 -0.8298821 -0.8275232 -0.8252065 -0.8247438 -0.8243418 -0.8243024 -0.8224756 -0.8215502 -0.8169271 -0.8163443 -0.808677	calreticulin eyes absent (Drosophila) homolog 2 ESTs ESTs period (Drosophila) homolog 1 ESTs interleukin 7 receptor DKFZP586A011 protein cortic al thymocyte receptor (X. laevis CTX) like serum amyloid A1 ESTs ESTs tyrosine 3-monooxygenase/tryptophan 5-monooxygenase activation protein, eta polypeptide

1555924 416434 322561 132711 70245	-0.8025163 -0.8006527 -0.7970805 -0.7962294 -0.795375	four and a half LIM domains 3 CSR1 protein SCN Circadian Oscillatory Protein (SCOP) ribosomal protein L31 Kruppel-like factor 5 (intestinal) Homo sapiens mRNA full length insert cDNA clone EUROIMAGE 50374 ESTs, Moderately similar to LONG-CHAIN FATTY ACID TRANSPORT PROTEIN [M.musculus]
250883 346545 1878409 530958 40027	-0.7887371 -0.7869766 -0.7809384 -0.7779473	ubiquitin-activating enzyme E1-like laminin, beta 1 catechol-O-methyltransferase smoothened (Drosophila) homolog
1474337 85840 290378	-0.7766266 -0.7754435 -0.7751825 -0.7736287	spinal cord-derived growth factor-B phosphorylase, glycogen; brain nicotinamide N-methyltransferase podocalyxin-like kallikrein 8 (neuropsin/ovasin) ATP-binding cassette, sub-family B (MDR/TAP), member 1
491403	-0.7700599	hypothetical protein FLJ10875 tumor necrosis factor receptor superfamily, member 1B
681992		GTP-binding protein overexpressed in skeletal muscle Homo sapiens cDNA FLJ13384 fis, clone PLACE1001062, highly similar to Homo sapiens mRNA for lysine-ketoglutarate reductase/saccharopine dehydrogenase
139660 869450 162308 209537	-0.7677153 -0.7659922 -0.7654139 -0.763749	RalGDS-like gene ESTs ribosomal protein L11 zinc finger protein 221 KIAA0603 gene product

810728 366591		hypothetical gene ZD52F10 T-cell lymphoma invasion and metastasis 2
32489		collagen, type XVI, alpha 1 hypothetical protein DKFZp566A1524
		synaptogyrin 1 receptor tyrosine kinase-like orphan receptor 1
		dystrobrevin, beta placental growth factor, vascular endothelial growth factor-related protein
188335	-0.7399294	egf-like module containing, mucin-like, hormone receptor-like sequence 2
	-0.7380353 -0.7364434	villin 1 Homo sapiens mRNA; cDNA DKFZp586B211 (from clone DKFZp586B211)
	-0.736252 -0.7346408	keratin 15 secretory leukocyte protease inhibitor (antileukoproteinase)
1630990 280907 742562 73252	-0.7307519 -0.730047	ribosomal protein L29 Kruppel-type zinc finger protein chromosome 16 open reading frame 5 3-hydroxy-3-methylglutaryl-Coenzyme A synthase 1 (soluble)
26566 1471829		protein-O-mannosyltransferase 1 uncharacterized hypothalamus protein HSMNP1
289428	-0.7270494	neurotrophic tyrosine kinase, receptor, type 2
898092 34150 24958	-0.7263513	connective tissue growth factor ESTs Homo sapiens mRNA; cDNA DKFZp434C2016 (from clone DKFZp434C2016)
725680	-0.7197488	transcription factor AP-2 gamma (activating enhancer-binding protein 2 gamma)
1686766	-0.7192745	complement component 3 Rag D protein integral membrane protein 3

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431231	-0.7182897	EGF-containing fibulin-like extracellular matrix protein 2
1908834	-0.7177051	ubiquitin A-52 residue ribosomal protein fusion product 1
207735	-0.7167416	serine (or cysteine) proteinase inhibitor, clade A (alpha-1 antiproteinase, antitrypsin), member 1
754157	-0.7165332	Homo sapiens mRNA; cDNA DKFZp434K2172 (from clone DKFZp434K2172)
35828	-0.7163689	diphtheria toxin receptor (heparin-binding epidermal growth factor-like growth factor)
1946448	-0.7150027	caveolin 2
470148	-0.712917	ESTs
840944	-0.7099353	early growth response 1
809784	-0.7095414	kallikrein 6 (neurosin, zyme)
877827		ribosomal protein S27a
32493	-0.7093777	integrin, alpha 6
813520	-0.7092422	EphB3
782501	-0.7086066	hypothetical protein PP1665
27544	-0.708383	prominin (mouse)-like 1
184022	-0.7070266	amyloid beta (A4) precursor protein-binding, family B, member 1 (Fe65)
323238	-0.7042807	GRO1 oncogene (melanoma growth stimulating activity, alpha)
488956	-0.70192	CUG triplet repeat, RNA-binding protein 2
1926246		ESTs, Moderately similar to T46371 hypothetical protein DKFZp434P1018.1 [H.sapiens]
325365	-0.7012106	HIV-1 rev binding protein 2
		checkpoint suppressor 1
		heat shock 27kD protein 2
		KIAA1183 protein
503335		hypothetical protein FLJ11196
757873		cyclin-dependent kinase 5, regulatory subunit 1 (p35)
		A kinase (PRKA) anchor protein (yotiao) 9 E74-like factor 5 (ets domain transcription factor)
		nucleoporin 88kD KIAA0172 protein

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46129	-0.6919169	ESTs, Highly similar to T17245 hypothetical protein DKFZp586J0917.1 [H.sapiens]
270917 789382 611481 35300 50892	-0.6898974 -0.6894079 -0.688788	secreted frizzled-related protein 1 Notch (Drosophila) homolog 4 HMG-box transcription factor TCF-3 KIAA0869 protein Homo sapiens, clone MGC:9913 IMAGE:3870821, mRNA,
		complete cds matrilin 2 KIAA1404 protein cholinergic receptor, nicotinic, beta polypeptide 1 (muscle)
		ribosomal protein L34 c-fos induced growth factor (vascular endothelial growth factor D)
		integrin, alpha 10 microtubule-associated protein, RP/EB family, member 2
839796	-0.6807912	candidate tumor suppressor p33 ING1 homolog
726582 50586 491763 52096	-0.6794282 -0.679034	actin-related protein 3-beta KIAA1545 protein interleukin 1, beta platelet-derived growth factor receptor, alpha polypeptide
		collagen, type VI, alpha 1 alcohol dehydrogenase 1C (class I), gamma polypeptide
41869	-0.6747514 -0.6740833	ribosomal protein S25 hypothetical protein FLJ11017 KIAA0015 gene product Homo sapiens clone HH409 unknown mRNA
773373 308539	-0.6730112 -0.6726187	hypothetical protein MGC14258 Homo sapiens cDNA FLJ12777 fis, clone NT2RP2001720
343760	-0.6724144	SH3 domain binding glutamic acid-rich protein like 2
		collagen, type IV, alpha 2 enhancer of filamentation 1 (cas-like docking; Crk-associated substrate related)
70201 sd-71385	-0.6675012	mitochondrial solute carrier

897731 296880	-0.6670507 -0.6655878	inositol 1,4,5-triphosphate receptor, type 2 latrophilin membrane protein, palmitoylated 1 (55kD) heparin-binding growth factor binding protein
	-0.6624986 -0.6591675 -0.6585351 -0.65723	hypothetical protein FLJ21841 hypothetical protein DKFZp762A227 lipoma HMGIC fusion partner KIAA0440 protein proline arginine-rich end leucine-rich repeat protein
744918	-0.6552361	Homo sapiens mRNA; cDNA DKFZp761I0911 (from clone DKFZp761I0911)
39600 795288		adenylate kinase 5 ubiquitin specific protease 4 (proto-oncogene)
1665444 774078 415233 307029 78946	-0.6508021 -0.6506593 -0.6500872	tumor endothelial marker 1 precursor leiomodin 1 (smooth muscle) ribosomal protein L37a ribosomal protein L26 Homo sapiens pyruvate dehydrogenase kinase 4 mRNA, 3' untranslated region, partial sequence
785733 415415		hypothetical protein FLJ12892 EST, Moderately similar to RL1X_HUMAN 60S RIBOSOMAL PROTEIN L18A [H.sapiens]
858167	-0.6453859	fatty-acid-Coenzyme A ligase, long-chain 4
823714	-0.642903	nuclear receptor co-repressor/HDAC3 complex subunit
193913	-0.6425581	v-yes-1 Yamaguchi sarcoma viral related oncogene homolog
220395	-0.6418175	hypothetical protein FLJ23293 similar to ARL-6 interacting
840942	-0.6383764	protein-2 major histocompatibility complex, class II, DP beta 1
1492147	-0.6371261 -0.636656 -0.6364598 -0.6362307	ribosomal protein S4, X-linked ESTs
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796152	-0.6356106	Homo sapiens cDNA FLJ11685 fis, clone HEMBA1004934
190059	-0.634814	guanine nucleotide binding protein (G protein), gamma 7
840511 81409	-0.6341661 -0.6338327	vimentin GABA(A) receptor-associated protein like 1
530036	-0.6324794	hypothetical protein FLJ20607 hypothetical protein FLJ13078 solute carrier family 25 (mitochondrial carrier; adenine nucleotide translocator), member 6
785538	-0.6289951	Homo sapiens mRNA; cDNA DKFZp434N2116 (from clone DKFZp434N2116)
		secretory carrier membrane protein 1
	-0.6263561 -0.6253276	fibroblast growth factor receptor 1 (fms-related tyrosine kinase 2, Pfeiffer syndrome)
774754	-0.6243946	catenin (cadherin-associated protein), beta 1 (88kD)
1572298	-0.6228686	CD3Z antigen, zeta polypeptide (TiT3 complex)
		ribosomal protein S10
		inositol polyphosphate-1-phosphatase
		Cdc42 effector protein 3
		epidermal growth factor (beta-urogastrone)
241489		adrenergic, beta-2-, receptor, surface
		ras homolog gene family, member C
	-0.6204409	
150897		hypothetical protein FLJ20241 UDP-GlcNAc:betaGal beta-1,3-N-
130091	-0.0100204	acetylglucosaminyltransferase 3
823851	-0.6182341	AE-binding protein 1
810057		vasoactive intestinal peptide receptor 1
	-0.6176265	· ·
		frizzled (Drosophila) homolog 1
		KIAA0300 protein
		BCE-1 protein
250797	-0.6124667	hypothetical protein FLJ20038
754040	-0.610807	NS1-associated protein 1
1584628 sd-71385	-0.6102937	alpha-actinin-2-associated LIM protein

sd-71385

	-0.6054867	hypothetical protein FLJ10390 ART-4 protein erythrocyte membrane protein band 4.1-like 2
342349	-0.603487	mitogen-activated protein kinase kinase kinase 14
67067	-0.6003961	Homo sapiens mRNA; cDNA DKFZp564O222 (from clone DKFZp564O222)
1533710	-0.5988282	ortholog of mouse integral membrane glycoprotein LIG-1
1707637 1634832 1609625	-0.5964339 -0.5931096 -0.5920312	hypothetical protein ESTs class I cytokine receptor selectin P ligand mammalian inositol hexakisphosphate kinase 2
897720 1472735	-0.5904719 -0.5888169	alpha2,3-sialyltransferase trophinin metallothionein 1E (functional) mitogen-activated protein kinase 8 interacting protein 2
1651906 564801		hypothetical protein FLJ23544 Homo sapiens cDNA: FLJ21409 fis, clone COL03924
248886	-0.5823315	rab3 GTPase-activating protein, non-catalytic subunit (150kD)
250654	-0.5815576	secreted protein, acidic, cysteine-rich (osteonectin)
1968422	-0.5802058	Homo sapiens mRNA full length insert cDNA clone EUROIMAGE 1968422
		LIM domain only 2 (rhombotin-like 1) Homo sapiens mRNA for FLJ00074 protein, partial cds
346902	-0.5794362	ESTs, Weakly similar to ALU2_HUMAN ALU SUBFAMILY SB SEQUENCE CONTAMINATION WARNING ENTRY [H.sapiens]
782193 358531	-0.5788147 -0.5784457	thioredoxin v-jun avian sarcoma virus 17 oncogene homolog
2271240	-0.5782432	hypothetical protein

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	120106	-0.5770131 -0.5768989	endothelin 3 caspase 1, apoptosis-related cysteine protease (interleukin 1, beta, convertase) B-cell associated protein ESTs, Weakly similar to ALU1_HUMAN ALU SUBFAMILY J SEQUENCE CONTAMINATION WARNING ENTRY [H.sapiens]
			gycosyltransferase laminin, gamma 2 (nicein (100kD), kalinin (105kD), BM600 (100kD), Herlitz junctional epidermolysis bullosa))
The second secon	489755	-0.5730453	a disintegrin and metalloproteinase domain 12 (meltrin alpha)
**************************************			angiopoietin 2 Homo sapiens cDNA: FLJ22290 fis, clone HRC04405
	••••		CDC-like kinase 3 ESTs, Highly similar to T12495 hypothetical protein DKFZp434H071.1 [H.sapiens]
			frizzled-related protein peroxisomal membrane protein 3 (35kD, Zellweger syndrome)
	1404774 814815 462939	-0.5686324 -0.5685596 -0.5675649 -0.5662456 -0.5660209	
	1636360	-0.5629882	DKFZP727C091 protein hypothetical protein FLJ14957 ESTs, Moderately similar to MAS2_HUMAN MANNAN- BINDING LECTIN SERINE PROTEASE 2 PRECURSOR [H.sapiens]
		-0.5618289	inositol polyphosphate phosphatase-like 1 ESTs ubiquitin-like 1 (sentrin)

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491519	-0.561401	Homo sapiens clone 24775 mRNA sequence
377314	-0.561312	casein kinase 2, alpha prime polypeptide
208969	-0.561267	FST
		KIAA1479 protein
		•
		integrin, alpha 1
14/3131	-0.5594101	transducin-like enhancer of split 2, homolog of Drosophila
		E(sp1)
812959	-0.5591846	KIAA1638 protein
868169	-0.5584677	lipoprotein lipase
		ribosomal protein, large, P0
		aryl hydrocarbon receptor nuclear translocator-like
30313	-0.5577701	ary riyurocarbon receptor nuolear translocator into
299720	-0.5565137	
	-0.555725	vasoactive intestinal peptide receptor 1
51981		ribosomal protein L7a
		KIAA1858 protein
		•
238907	-0.5534598	hypothetical protein, clone
		Telethon(Italy_B41)_Strait02270_FL142
		DHHC1 protein
282404	-0.5528532	Homo sapiens mRNA for KIAA1671 protein, partial cds
044500	0.55446	Itallikasia E
	-0.55146	
746373	-0.5513143	RNA polymerase I transcription factor RRN3
491186	-0.5507413	Homo sapiens cDNA: FLJ23131 fis, clone LNG08502
		poly(A)-binding protein, cytoplasmic 1-like
345764	-0.5504113	special AT-rich sequence binding protein 1 (binds to nuclear
		matrix/scaffold-associating DNA's)
4.533000	0.5400044	FOT
	-0.5496944	
288748	-0.5493871	Homo sapiens, clone MGC:5352 IMAGE:3048106, mRNA,
		complete cds
810097	-0.5487174	Homo sapiens cDNA: FLJ21721 fis, clone COLF0381
		•
755881	-0.5473964	aquaporin 5
1568391	-0.5467858	plastin 3 (T isoform)
		v-raf-1 murine leukemia viral oncogene homolog 1
20000	0.0407010	V fat i filatillo loakonna vital onoogono nomolog .
503671	-0.5461871	Homo sapiens cDNA FLJ14368 fis, clone HEMBA1001122
		•
2388571	-0.5459823	A kinase (PRKA) anchor protein 8
2388571 sd-71385	-0.5459823	A kinase (PRKA) anchor protein 8

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		cellular retinoic acid-binding protein 1 retinoic acid receptor responder (tazarotene induced) 2
	-0.5446617 -0.5443525	ESTs Homo sapiens cDNA FLJ10205 fis, clone HEMBA1004954
2213824 1597813	-0.5412089 -0.5406089	hypothetical protein FLJ12806 protease inhibitor 3, skin-derived (SKALP) hypothetical protein PP1044 acyl-Coenzyme A dehydrogenase, very long chain
341759	-0.5403406	lung type-I cell membrane-associated glycoprotein
42681	-0.5388424	ribosomal protein S4, Y-linked NY-REN-25 antigen eukaryotic translation elongation factor 1 alpha 1
162775	-0.5385174	DEAD/H (Asp-Glu-Ala-Asp/His) box polypeptide 5 (RNA
	-0.5382125	helicase, 68kD) TBP-associated factor 172 ribosomal protein L35 eukaryotic translation elongation factor 1 alpha 1
	-0.5376323 -0.5373697	ESTs Homo sapiens mRNA; cDNA DKFZp586N012 (from clone DKFZp586N012)
233679		ESTs hypothetical protein FLJ22362 sushi-repeat-containing protein, X chromosome
825356	-0.533771	Homo sapiens cDNA FLJ11997 fis, clone HEMBB1001458
767982	-0.533231	Homo sapiens, clone IMAGE:4134852, mRNA, partial cds
277627	-0.5329107	Human SH3 domain-containing protein SH3P18 mRNA,
346688 712023 24729 1590021	-0.5315035 -0.5302372 -0.5302207	complete cds KIAA0819 protein melanoma inhibitory activity AT-binding transcription factor 1 cholinergic receptor, muscarinic 1 ets variant gene 6 (TEL oncogene)

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811779	-0.5291718	phosphoinositide-3-kinase, regulatory subunit, polypeptide 1 (p85 alpha)
488130	-0.5289721	Homo sapiens cDNA FLJ20767 fis, clone COL06986
159462 1570427 2308346 1881774	-0.5280955 -0.5268833 -0.5265891	serum constituent protein hypothetical protein MGC4309 cyclin-dependent kinase 2 KIAA1678 protein kinase C and casein kinase substrate in neurons 2
	-0.5253542 -0.5253354	myosin IE Rac/Cdc42 guanine exchange factor (GEF) 6
753620 80374		insulin-like growth factor binding protein 6 pyruvate dehydrogenase (lipoamide) alpha 1
		transgelin 2 Homo sapiens mRNA; cDNA DKFZp761P0114 (from clone DKFZp761P0114)
855061 183704 155896	-0.5203046 -0.5184644 -0.5182247	PRO1659 protein vascular endothelial growth factor B Homo sapiens, clone MGC:13446 IMAGE:4275731, mRNA, complete cds ORF adenosine deaminase, RNA-specific, B1 (homolog of rat RED1)
685185 753301 795178 1911663	-0.5172158 -0.5171758 -0.5169102 -0.5168962	reticulon 2 carcinoembryonic antigen-related cell adhesion molecule 1 (biliary glycoprotein) lactate dehydrogenase C
868308 809998 771004 842896	-0.5157133 -0.5144681 -0.5144276 -0.5137089	peptidylprolyl isomerase E (cyclophilin E) ribosomal protein S23 amylase, alpha 2A; pancreatic KIAA1201 protein hypothetical protein DKFZp762L0311 tumor necrosis factor receptor superfamily, member 6

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341763	-0.4953701	caspase 5, apoptosis-related cysteine protease
1/17886	-0.4948424 -0.4941304 -0.4934665	ESTs hypothetical protein FLJ23239 Homo sapiens cDNA: FLJ22130 fis, clone HEP19632
868400 809894 753745	0.401862	glutaminyl-tRNA synthetase acetyl-CoA synthetase ESTs, Weakly similar to S57447 HPBRII-7 protein [H.sapiens]
813698 253884 131012 855755	-0.4908497 -0.4898644 -0.4897811 -0.4897744 -0.4897236	sprouty (Drosophila) homolog 2 Human BAC clone GS1-99H8 hypothetical protein FLJ10633
2413337	-0.4879268	sortilin-related receptor, L(DLR class) A repeats-containing
327432 1897944	-0.4870045 -0.4855156	semaphorin Y ESTs, Weakly similar to T00366 hypothetical protein KIAA0669 [H.sapiens]

Example VI: Genes for discriminating between normal and ADH (non-malignant) versus DCIS and IDC (malignant)

As shown in Table 5 below, 400 genes were identified as being able to discriminate between normal and ADH (non-malignant) versus DCIS and IDC (malignant).

Table 5

10010		
ClonelD 796469	Weight 1.4207633	Description HSPC150 protein similar to ubiquitin-conjugating
488964	1.3447179	enzyme H2A histone family, member O

sd-71385

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1505038 1500000 1554549 812238 35147 122077 788654 595037 565319	1.3277637 1.2926116 1.2787033 1.2664748 1.2581066 1.2576139 1.2573483 1.2486446 1.2155833	hypothetical protein FLJ20171 H2B histone family, member B hydroxyacyl glutathione hydrolase hypothetical protein MGC4692 ESTs, Weakly similar to unnamed protein product [H.sapiens] putative membrane protein growth factor receptor-bound protein 2 retinoic acid induced 3 Homo sapiens mRNA; cDNA DKFZp564B1264 (from clone DKFZp564B1264)
283919 1917941 359887 471568	1.2112507 1.1872008 1.1837896 1.1673113	H2A histone family, member L purine-rich element binding protein B translocase of inner mitochondrial membrane 17 (yeast) homolog A hematological and neurological expressed 1
290841 796694	1.1670252 1.1580364	H2B histone family, member A baculoviral IAP repeat-containing 5 (survivin)
366132	1.1440486	succinate dehydrogenase complex, subunit C, integral membrane protein, 15kD
823598 1323448	1.1425746 1.1309009	proteasome (prosome, macropain) 26S subunit, non-ATPase, 12 cysteine-rich protein 1 (intestinal)
810711	1.1074523	stearoyl-CoA desaturase (delta-9-desaturase)
741474 745606 2054635	1.0870449 1.0663136 1.0613961	glucose phosphate isomerase hypothetical protein PP591 proteasome (prosome, macropain) subunit, alpha
178805	1.0513873	type, 7 Human DNA sequence from clone RP5-850E9 on chromosome 20. Contains part of the gene for a novel C2H2 type zinc finger protein similar to Drosophila Scratch (Scrt), Slug and Xenopus Snail, a novel gene similar to Drosophila CG6762, STSs, GSSs and five CpG
347373	1.0454939	transcription elongation factor B (SIII), polypeptide 1 (15kD, elongin C)

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199403	1.0142329	lectin, galactoside-binding, soluble, 8 (galectin 8)
814054 2029173	1.0130821 1.0094499	KIAA0040 gene product ESTs, Weakly similar to N-WASP [H.sapiens]
209066 782428 769921 470061 796723	1.0066096 1.0008279 0.9944462 0.9920108 0.9915885	KIAA0250 gene product ubiquitin carrier protein E2-C seven in absentia (Drosophila) homolog 2 Homo sapiens clone CDABP0014 mRNA sequence
1609836	0.9886168	glutamate-ammonia ligase (glutamine synthase)
2322367 280375 2016908	0.9859632 0.9793036 0.969649	reticulon 4 PRO2000 protein ESTs, Weakly similar to CA13_HUMAN COLLAGEN ALPHA 1(III) CHAIN PRECURSOR [H.sapiens]
1858892 46248	0.9669022 0.9628117	hypothetical protein MGC4825 ADP-ribosyltransferase (NAD+; poly (ADP-ribose) polymerase)
2014034	0.9619198	methylene tetrahydrofolate dehydrogenase (NAD+ dependent), methenyltetrahydrofolate cyclohydrolase
898032	0.9547022	KIAA0097 gene product
725454	0.9476507	CDC28 protein kinase 2
79520	0.9457391	RAB2, member RAS oncogene family
810558	0.943041	proteasome (prosome, macropain) 26S subunit, ATPase, 4
272529	0.9423688	phosphomannomutase 2
122241	0.939204	proteasome (prosome, macropain) subunit, beta type, 2
469686	0.9381847	Ric (Drosophila)-like, expressed in many tissues
624667 488202	0.9336899 0.932881	CGI-92 protein ESTs, Weakly similar to YZ28_HUMAN HYPOTHETICAL PROTEIN ZAP128 [H.sapiens]
825470 1640821	0.9274271 0.920874	topoisomerase (DNA) II alpha (170kD) ESTs, Weakly similar to I78885 serine/threonine- specific protein kinase [H.sapiens]

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686552 2016648	0.9182272 0.917373	golgi phosphoprotein 1 Homo sapiens mRNA; cDNA DKFZp434N1728 (from clone DKFZp434N1728)
1911343 781097 244801	0.9110591 0.9100493 0.908481	RAB26, member RAS oncogene family reticulon 3 Rho guanine exchange factor (GEF) 11
754628 1574058	0.9061145 0.9058211	ESTs 1-acylglycerol-3-phosphate O-acyltransferase 2 (lysophosphatidic acid acyltransferase, beta)
753299	0.9019921	hypothetical protein FLJ10504
811774	0.9014801	CGI-49 protein
595213	0.8978322	hypothetical protein
868128	0.8970146	JM4 protein
810124	0.8891272	platelet-activating factor acetylhydrolase, isoform
00400	0.0000000	lb, gamma subunit (29kD) hypothetical protein DKFZp762E1312
66406	0.8888639	hypothetical protein DNF2p702E1312
1636092	0.8841788	hypothetical protein FLJ20657 hypothetical protein MGC2745
1869201	0.8825788	phosphoenolpyruvate carboxykinase 2
625923	0.8821749	(mitochondrial)
1492238	0.8804306	HSPC003 protein
731044	0.8716644	glutaredoxin 2
839682	0.8696528	ubiquitin-conjugating enzyme E2N (homologous to yeast UBC13)
951233	0.8614727	proteasome (prosome, macropain) subunit, beta type, 3
843195	0.8606568	phosphoserine phosphatase
288999	0.8593924	small protein effector 1 of Cdc42
51773	0.8588635	hypothetical protein MGC3077
209066	0.8582298	serine/threonine kinase 15
1474955	0.8573467	TATA box binding protein (TBP)-associated factor, RNA polymerase II, N, 68kD (RNA-binding protein 56)
2043167	0.8551193	BCL2-associated athanogene 3
742707	0.8515067	ESTs, Weakly similar to MUC2_HUMAN MUCIN 2 PRECURSOR [H.sapiens]
743589	0.8514377	ESTs, Weakly similar to T2D3_HUMAN TRANSCRIPTION INITIATION FACTOR TFIID 135 KDA SUBUNIT [H.sapiens]

704414	0.8498631	small nuclear ribonucleoprotein polypeptides B and B1
2309073	0.8484971	frizzled (Drosophila) homolog 5
2052113	0.8477245	hypothetical protein FLJ10903
686172	0.846207	KIAA0008 gene product
150003	0.8447372	hypothetical protein FLJ13187
705064	0.8401441	transforming, acidic coiled-coil containing protein
1709791	0.8397779	BAI1-associated protein 1
1469425	0.8391993	SRY (sex determining region Y)-box 22
429799	0.8386406	hypothetical protein FLJ21939 similar to 5- azacytidine induced gene 2
729975	0.8385313	meningioma expressed antigen 5 (hyaluronidase)
1393018	0.8376676	general transcription factor IIIC, polypeptide 1 (alpha subunit, 220kD)
1492463	0.8360771	selenoprotein X, 1
2028949	0.8358024	hypothetical protein PRO1855
789012	0.8351735	fibulin 2
470124	0.8347241	RAD1 (S. pombe) homolog
1409509	0.8339967	troponin T1, skeletal, slow
1605426	0.8317254	hypothetical protein FLJ13352
44292	0.831599	Homo sapiens mRNA; cDNA DKFZp434C107 (from clone DKFZp434C107)
1435003	0.8311727	tumor suppressing subtransferable candidate 1
503215	0.8286483	pilin-like transcription factor
504308	0.8199799	hypothetical protein FLJ10540
785707	0.8176557	protein regulator of cytokinesis 1
1500162	0.8155519	ESTs
149355	0.8134342	translocating chain-associating membrane protein
1845169	0.8131362	RAB35, member RAS oncogene family
869375	0.808792	isocitrate dehydrogenase 2 (NADP+), mitochondrial
1492426	0.8032542	chromosome 19 open reading frame 3
782513	0.8030042	interferon, alpha-inducible protein (clone IFI-6-16)
813281	0.8016742	WW domain-containing protein 1
814378	0.8015236	serine protease inhibitor, Kunitz type, 2

700792	0.8005506	cyclin-dependent kinase inhibitor 3 (CDK2-associated dual specificity phosphatase)
340558	0.7990715	actin related protein 2/3 complex, subunit 5 (16 kD)
842994	0.7983201	cathepsin Z
138189	0.7953361	Wolfram syndrome 1 (wolframin)
289978	0.7931469	ubiquitin-like 4
2019223	0.7893602	mitochondrial ribosomal protein L17
2110511	0.7863117	artemin
814528	0.7845635	Homo sapiens cDNA: FLJ22139 fis, clone HEP20959
1573251	0.7816689	peroxisomal long-chain acyl-coA thioesterase
773922	0.7799164	KIAA0005 gene product
172517	0.7779159	hippocalcin-like 1
564981	0.7745626	ESTs
1420370	0.7726877	biliverdin reductase B (flavin reductase (NADPH))
308466	0.771216	GTP-binding protein Sara
199645	0.769591	nicastrin
1422338	0.7690604	ribonucleotide reductase M2 polypeptide
1474424	0.7689082	Homo sapiens cDNA FLJ12758 fis, clone NT2RP2001328
813751	0.7665427	sialyltransferase 4C (beta-galactosidase alpha- 2,3-sialytransferase)
131094	0.7626373	Homo sapiens cDNÁ: FLJ21587 fis, clone COL06946
842980	0.762412	developmentally regulated GTP-binding protein 1
212542	0.7585053	Homo sapiens cDNA FLJ12900 fis, clone NT2RP2004321
108425	0.7581954	
811761	0.7578696	Nijmegen breakage syndrome 1 (nibrin)
241348	0.757138	prenylcysteine lyase
810725	0.7561061	ATPase, H+ transporting, lysosomal (vacuolar proton pump) 21kD
1460110	0.7559869	proteasome (prosome, macropain) subunit, beta type, 5
120749	0.7545697	ESTs, Moderately similar to KIAA1215 protein [H.sapiens]
827171	0.7543905	ESTs

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39884	0.7526496	IMP (inosine monophosphate) dehydrogenase 1
124298	0.7507816	microsomal glutathione S-transferase 3
753378	0.7492212	hypothetical protein FLJ22649 similar to signal peptidase SPC22/23
210862	0.7483198	acyl-Coenzyme A oxidase 1, palmitoyl
785766	0.7476331	hypothetical protein
1553306	0.7471557	proteasome (prosome, macropain) 26S subunit, non-ATPase, 11
470099	0.7443747	HT002 protein; hypertension-related calcium- regulated gene
83363	0.7435058	protein-L-isoaspartate (D-aspartate) O- methyltransferase
489351	0.7422879	hypothetical protein DKFZp566J2046
810612	0.7419597	S100 calcium-binding protein A11 (calgizzarin)
825585	0.741258	tubulin-specific chaperone e
1456348	0.7401571	N-acetylneuraminic acid phosphate synthase; sialic acid synthase
1473922	0.7399948	actin related protein 2/3 complex, subunit 3 (21 kD)
150314	0.739762	lysophospholipase l
897806	0.7388301	hypoxia-inducible factor 1, alpha subunit (basic helix-loop-helix transcription factor)
811585	0.738212	huntingtin (Huntington disease)
685516	0.7378926	putative G protein-coupled receptor
76605	0.7376829	nesca protein
1476053	0.7367106	RAD51 (S. cerevisiae) homolog (E coli RecA homolog)
221295	0.7342275	inhibitor of DNA binding 2, dominant negative helix-loop-helix protein
824879	0.7333071	hypothetical protein MGC11275
768570	0.732628	hypothetical protein FLJ11280
73009	0.731401	ESTs, Weakly similar to A43932 mucin 2
EC4400	0.7004005	precursor, intestinal [H.sapiens]
564492 2017415	0.7301895	mitochondrial carrier homolog 2
2017415	0.7281714	centromere protein A (17kD)
788654 488505	0.7244749	accessory proteins DAD24/DAD20
488505 824962	0.7242619 0.7238234	accessory proteins BAP31/BAP29 karyopherin alpha 2 (RAG cohort 1, importin alpha 1)

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1616253 1435862	0.7231756 0.723081	breast carcinoma amplified sequence 1 antigen identified by monoclonal antibodies 12E7, F21 and O13
786067	0.7228184	cell division cycle 25B
2050827	0.7211304	proteasome (prosome, macropain) 26S subunit, ATPase, 5
248649	0.7210707	hypothetical protein FLJ13910
51532	0.7195798	ADP-ribosylation factor-like 6 interacting protein
727078	0.7184618	Homo sapiens cDNA: FLJ23602 fis, clone LNG15735
37708	0.7163849	hypothetical protein MGC3101
430235	0.7162503	H2B histone family, member Q
897770	0.715754	
292936	0.7154295	hypothetical protein FLJ10468
365738	0.7152855	ESTs
845363	0.714201	non-metastatic cells 1, protein (NM23A) expressed in
809944	0.7139515	KIAA0310 gene product
1631699	0.7115561	valosin-containing protein
813629	0.7102574	YME1 (S.cerevisiae)-like 1
813410	0.7093173	polymerase (RNA) II (DNA directed) polypeptide K (7.0kD)
826256	0.7092586	transmembrane 7 superfamily member 1 (upregulated in kidney)
124331	0.7081481	cleavage and polyadenylation specific factor 5, 25 kD subunit
770845	0.7068283	hexokinase 1
67765	0.7064266	carboxypeptidase M
207288	0.7058528	insulin induced gene 1
1639531	0.7033264	RAB27A, member RAS oncogene family
731023	0.7032815	WD repeat domain 5
756442	0.7016064	P450 (cytochrome) oxidoreductase
358162	0.6999211	protein predicted by clone 23627
782608	0.698569	mitochondrial ribosomal protein L9
509823	0.6982362	carcinoembryonic antigen-related cell adhesion molecule 6 (non-specific cross reacting antigen)
345787	0.6981453	highly expressed in cancer, rich in leucine heptad repeats
810402	0.6978141	hypothetical protein

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744417	0.6975761	carnitine acetyltransferase
814306	0.6963874	tumor protein D52
41356	0.6961169	protein phosphatase 2, regulatory subunit B (B56), alpha isoform
813419	0.6951349	hydroxyacyl-Coenzyme A dehydrogenase, type II
629944	0.6950339	myosin VB
327506	0.6943362	Homo sapiens mRNA full length insert cDNA clone EUROIMAGE 327506
768064	0.6936336	cytochrome P450, subfamily I (aromatic compound-inducible), polypeptide 1
770992	0.693225	
469383	0.6925975	chromosome 8 open reading frame 1
1492780	0.6923749	Homo sapiens cDNA FLJ14459 fis, clone HEMBB1002409
41569	0.691145	hypothetical protein FLJ12650
509588	0.6903363	TATA box binding protein (TBP)-associated factor, RNA polymerase II, J, 20kD
144880	0.6902434	hypothetical protein from EUROIMAGE 1759349
789376	0.6883473	thioredoxin reductase 1
268946	0.6881593	WD40 protein Ciao1
810156	0.6879064	deoxythymidylate kinase (thymidylate kinase)
84295	0.6876167	interleukin 1 receptor antagonist
246800	0.6867481	hypothetical protein FLJ10803
589232	0.6865999	hypothetical protein FLJ11506
859761	0.6846134	poliovirus receptor-related 2 (herpesvirus entry mediator B)
431505	0.6840493	ESTs, Highly similar to A31026 probable membrane receptor protein [H.sapiens]
109863	0.683291	epithelial membrane protein 2
770355	0.6829507	lanosterol synthase (2,3-oxidosqualene- lanosterol cyclase)
344091	0.6818771	ESTs
813707	0.6814476	regulator of G-protein signalling 16
124781	0.6809199	squalene epoxidase
502774	0.6807524	hypothetical protein FLJ20623
825740	0.6805157	DKFZp434J1813 protein
1536006	0.6805004	ESTs

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203003	0.6802818	non-metastatic cells 4, protein expressed in
785616	0.6800658	signal sequence receptor, alpha (translocon- associated protein alpha)
785840	0.6800437	SEC24 (S. cerevisiae) related gene family, member D
256619	0.6797698	hydroxysteroid (17-beta) dehydrogenase 7
2108077	0.6796871	CGI-112 protein
742595	0.6784804	cyclin-dependent kinase 5
1476065	0.6783519	stathmin 1/oncoprotein 18
68636	0.6778817	hypothetical protein MGC2477
564847	0.6770455	ESTs
744047	0.6748419	polo (Drosophia)-like kinase
897813	0.6741253	polyadenylate binding protein-interacting protein
823930	0.6738266	actin related protein 2/3 complex, subunit 1A (41 kD)
236034	0.673506	uncoupling protein 2 (mitochondrial, proton carrier)
743977	0.6729967	Homo sapiens mRNA for TL132
293727	0.6728542	hypothetical protein MGC861
590759	0.672734	sterol-C4-methyl oxidase-like
811024	0.6706275	bone marrow stromal cell antigen 2
785795	0.6683919	hypothetical protein FLJ12910
530310	0.6682042	KIAA0143 protein
1734309	0.666164	sperm associated antigen 4
1898619	0.6651268	hypothetical protein MGC15737
358267	0.6648183	EST, Moderately similar to AF119917 63 PRO2831 [H.sapiens]
753400	0.6633015	CGI-204 protein
292770	0.6609782	Homo sapiens, clone IMAGE:3627860, mRNA, partial cds
884425	0.6608337	chaperonin containing TCP1, subunit 5 (epsilon)
1696757	0.6599512	hypothetical protein KIAA1165
121251	0.6596925	hypothetical protein MGC5576
770785	0.658751	1,2-alpha-mannosidase IC
590774	0.657447	mitogen-activated protein kinase 13
969877	0.6564571	synaptosomal-associated protein, 25kD
415102	0.6562436	cell division cycle 25C
120271	0.6562017	hypothetical protein MGC4692

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713782	0.6552297	a disintegrin and metalloproteinase domain 15 (metargidin)
49351	0.6551453	SEX gene
488642	0.6541915	ESTs, Weakly similar to I38022 hypothetical protein [H.sapiens]
343607	0.6510034	AD-015 protein
2306987	0.6501071	secreted and transmembrane 1
280249	0.64759	Kruppel-like factor 7 (ubiquitous)
752631	0.6473517	fibroblast growth factor receptor 3 (achondroplasia, thanatophoric dwarfism)
489594	0.645574	hypothetical protein FLJ11565
773188	0.6448143	nuclear receptor subfamily 1, group D, member 2
2015517	0.6441737	hypothetical protein FLJ22237
343731	0.6424907	mita abandrial riba comal protein 140
491524	0.6424448	mitochondrial ribosomal protein L13
824524	0.6424419	UDP-galactose transporter related
593431	0.6422897	ESTs, Moderately similar to CEGT_HUMAN CERAMIDE GLUCOSYLTRANSFERASE [H.sapiens]
812994	0.6420817	retinoid X receptor, alpha
1631735	0.6404771	Homo sapiens, clone IMAGE:3604336, mRNA, partial cds
278531	0.6404612	cytochrome c oxidase subunit VIc
2302099	0.6386686	sialidase 3 (membrane sialidase)
454896	0.6384516	DnaJ (Hsp40) homolog, subfamily A, member 2
1456701	0.6383709	B-cell CLL/lymphoma 9
2055807	0.638184	protein kinase domains containing protein similar to phosphoprotein C8FW
1518402	0.6378481	KIAA1361 protein
810762	0.6371461	SNARE protein
124447	0.6363079	KIAA1184 protein
49273	0.635859	solute carrier family 27 (fatty acid transporter), member 4
365060	0.6350631	RAB11A, member RAS oncogene family
1591264	0.6337293	transaldolase 1
41698	0.6327738	progesterone binding protein
810316	0.6323393	very long-chain acyl-CoA synthetase; lipidosin
826363	0.6322533	lysophospholipase II

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2011515 770675 1461477	0.6319712 0.6315109 0.6300096	DKFZP586B0923 protein Homo sapiens cDNA: FLJ21323 fis, clone COL02374 Homo sapiens mRNA; cDNA DKFZp586l0324
20024	0.600067	(from clone DKFZp586l0324)
366834 1601947	0.629867 0.6297475	envoplakin cytochrome c oxidase subunit VIIa polypeptide 2 (liver)
510575	0.6295917	hypothetical protein FLJ22087
503851	0.6291771	nuclear receptor co-repressor/HDAC3 complex subunit
810063	0.6286184	growth factor, erv1 (S. cerevisiae)-like (augmenter of liver regeneration)
754653	0.6277098	cadherin, EGF LAG seven-pass G-type receptor 3, flamingo (Drosophila) homolog
1518890	0.6251884	metallothionein-like 5, testis-specific (tesmin)
784105	0.6244805	ESTs
205049	0.6231646	protein kinase H11; small stress protein-like protein HSP22
325606	0.6225147	hypothetical protein MGC14353
760299	-1.8441097	dickkopf (Xenopus laevis) homolog 3
200814	-1.8295958	membrane metallo-endopeptidase (neutral endopeptidase, enkephalinase, CALLA, CD10)
1882697	-1.715818	peanut (Drosophila)-like 2
344720	-1.6567437	glycophorin C (Gerbich blood group)
1161564	-1.5877154	desmuslin
45099	-1.566311	regucalcin (senescence marker protein-30)
75859	-1.5303427	N-myc downstream-regulated gene 2
811920	-1.5255258	interleukin 11 receptor, alpha
1569187	-1.4872982	heparan sulfate (glucosamine) 3-O- sulfotransferase 4
796542	-1.4697418	ets variant gene 5 (ets-related molecule)
767202	-1.4573536	latent transforming growth factor beta binding protein 2
285377	-1.4496786	pellino (Drosophila) homolog 2
300632	-1.449587	hypothetical protein FLJ21044 similar to Rbig1

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160192	-1.4231519	ESTs, Weakly similar to 2004399A chromosomal protein [H.sapiens]
611532 813265	-1.3689616 -1.3632094	troponin I, skeletal, fast Homo sapiens mRNA; cDNA DKFZp564H1916 (from clone DKFZp564H1916)
1469377 810002	-1.359583 -1.3553375	lipoma HMGIC fusion partner-like 2 Homo sapiens, clone MGC:19762 IMAGE:3636045, mRNA, complete cds
2056139 377275	-1.3506352 -1.3392122	LIM domain protein ataxia-telangiectasia group D-associated protein
298122 72778	-1.3364021 -1.3299927	frizzled (Drosophila) homolog 7 caspase 7, apoptosis-related cysteine protease
841308	-1.3246996	myosin, light polypeptide kinase
377461	-1.3214357	caveolin 1, caveolae protein, 22kD
1473274	-1.3200174	myosin regulatory light chain 2, smooth muscle isoform
1554167	-1.3141843	hypothetical protein FLJ14529
131839	-1.3048208	folate receptor 1 (adult)
67741	-1.3014364	PP2135 protein
132857	-1.280579	Homo sapiens mRNA; cDNA DKFZp586N1323 (from clone DKFZp586N1323)
345670	-1.2805684	ESTs, Moderately similar to I59348 CCAAT binding transcription factor CBF subunit C - rat [R.norvegicus]
781014	-1.2659158	suppression of tumorigenicity 5
289760	-1.2610055	ESTs, Highly similar to T00391 hypothetical protein KIAA0612 [H.sapiens]
742685	-1.248683	disabled (Drosophila) homolog 2 (mitogen- responsive phosphoprotein)
52419	-1.2447753	Friedreich ataxia region gene X123
838478	-1.2349342	neurocalcin delta
839736	-1.2337192	crystallin, alpha B
1558675	-1.2313679	SRY (sex determining region Y)-box 10
796181	-1.2276581	growth arrest-specific 6
878836	-1.2200837	secretory granule, neuroendocrine protein 1 (7B2 protein)
130201	-1.2181641	intercellular adhesion molecule 2
1917449	-1.217371	serum amyloid A4, constitutive
sd-71385		

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22917	-1.2164705	Homo sapiens mRNA; cDNA DKFZp761M0111 (from clone DKFZp761M0111)
1871116	-1.209237	Homo sapiens mRNA; cDNA DKFZp434C1714 (from clone DKFZp434C1714); partial cds
823871	-1.2090693	SPARC-like 1 (mast9, hevin)
811837	-1.2030392	
811848	-1.2011809	hypothetical protein
840266	-1.200789	Homo sapiens cDNA: FLJ22667 fis, clone HSI08385
753071	-1.2005994	Homo sapiens cDNA: FLJ22528 fis, clone HRC12825
2106144	-1.1719133	regulated in glioma
2504881	-1.1674204	signal transducer and activator of transcription 5A
811837	-1.157587	eukaryotic translation elongation factor 1 alpha 1
486683	-1.1548164	Homo sapiens mRNA; cDNA DKFZp564J0323 (from clone DKFZp564J0323)
712139	-1.1491129	ADP-ribosylation factor-like 7
196435	-1.1475545	ESTs
877621	-1.1422087	nGAP-like protein
811088	-1.140941	ephrin-B3
322561	-1.1270333	ribosomal protein L31
712401	-1.1218827	phosphoinositide-3-kinase, catalytic, delta polypeptide
788234	-1.1061158	inhibitor of DNA binding 4, dominant negative helix-loop-helix protein
140574	-1.103064	small inducible cytokine subfamily D (Cys-X3-Cys), member 1 (fractalkine, neurotactin)
490023	-1.1020527	hypothetical protein MGC2648
1584540	-1.0933558	Homo sapiens mRNA; cDNA DKFZp586M2022 (from clone DKFZp586M2022)
1556433 47043 303109 343760	-1.0887923 -1.0841886 -1.0807576 -1.0803279	GRO3 oncogene tensin purinergic receptor (family A group 5) SH3 domain binding glutamic acid-rich protein like 2

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248631	-1.0784377	aminomethyltransferase (glycine cleavage system protein T)
740620	-1.071175	tropomyosin 2 (beta)
529843	-1.0698218	ESTs, Moderately similar to JC5238 galactosylceramide-like protein, GCP [H.sapiens]
990881	-1.0640807	
80344	-1.0602825	interleukin 7 receptor
625399	-1.0594452	hypothetical protein similar to beta-transducin family
416676	-1.0568729	pellino (Drosophila) homolog 1
416959	-1.0566462	nuclear factor I/B
307029	-1.0497874	ribosomal protein L26
488404	-1.0484323	Homo sapiens clone TUA8 Cri-du-chat region mRNA
291478	-1.0400846	runt-related transcription factor 3
153760	-1.0340645	EphB1
119290	-1.025797	cortic al thymocyte receptor (X. laevis CTX) like
814443	-1.0250673	hypothetical protein MGC3232
757191	-1.0238476	ESTs
208718	-1.0229324	annexin A1
161456	-1.0220494	serum amyloid A1
1587710	-1.0209983	period (Drosophila) homolog 1
160609	-1.0208819	ESTs
593023	-1.0158099	dystrobrevin, beta
1878409	-1.013872	catechol-O-methyltransferase
781017	-1.0125987	early growth response 2 (Krox-20 (Drosophila) homolog)
753162	-1.0119485	KIAA0603 gene product
897963	-1.0075423	phosphatidic acid phosphatase type 2A
505864	-1.0033263	RalGDS-like gene
323780	-1.0027215	Homo sapiens cDNA FLJ11177 fis, clone PLACE1007402
343695	-1.002495	hypothetical protein FLJ10875
277571	-0.9977509	KIAA1706 protein
869450	-0.9934083	ribosomal protein L11
280907	-0.9932806	Kruppel-type zinc finger protein
290378	-0.9932643	podocalyxin-like
1635062	-0.9905245	DKFZP586A011 protein

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270826	-0.9896888	Homo sapiens cDNA FLJ13329 fis, clone
		OVARC1001795
1257131	-0.9869928	ESTs

Example VII: Genes for discriminating between ADH and DCIS

As shown in Table 6 below, 350 genes were identified as being able to discriminate between ADH and DCIS. The actual data is shown in Figure 3.

Table 6

ClonelD	Weight	Description
1404774	-0.5685596 parathyroid	l hormone-like hormone
823871	-0.9429443 SPARC-lik	e 1 (mast9, hevin)
1882697	-1.4318896 peanut (Dr	osophila)-like 2
140071	-0.5708303 frizzled-rela	ated protein
160192	-1.1751869 ESTs, Wea [H.sapiens	akly similar to 2004399A chromosomal protein
796542	-1.2625621 ets variant	gene 5 (ets-related molecule)
611532	-1.3158379 troponin I,	skeletal, fast
1473274	-1.1172693 myosin reg	ulatory light chain 2, smooth muscle isoform
469306	-0.5113377 gastrin-rele	easing peptide
2306697	-0.4494025 neuromedi	n B
132857	-1.0084069 Homo sapi DKFZp586	ens mRNA; cDNA DKFZp586N1323 (from clone N1323)
2504881	-1.000863 signal trans	sducer and activator of transcription 5A
760299	-1.7010462 dickkopf (X	enopus laevis) homolog 3
293819	-0.4051233 oxidoreduc	tase UCPA
130835	-	ens, Similar to clone FLB3816, clone 54380, mRNA
66532	-0.5773248 endothelin	3
2499829	•	protein 145 (Kruppel-like, expressed in rtic leukemia)
85840	-0.7754435 nicotinamic	•
sd-71385		

859359 200814	 -0.4669761 quinone oxidoreductase homolog -1.4325283 membrane metallo-endopeptidase (neutral endopeptidase, enkephalinase, CALLA, CD10)
825287	-0.2783909 tumor necrosis factor (ligand) superfamily, member 11
202577 2014373 153760 377275	-0.2283149 histamine N-methyltransferase 0.0232775 HNK-1 sulfotransferase -0.9227865 EphB1 -1.3102234 ataxia-telangiectasia group D-associated protein
745490 172783 1558108	-0.6329354 hypothetical protein FLJ20607 -0.6088873 hypothetical protein FLJ10390 0.0044098 ATP-binding cassette, sub-family C (CFTR/MRP), member 8
1587710 141731 490484 504959	-0.8252065 period (Drosophila) homolog 1 -0.4531468 -0.3388325 ESTs -0.1620065 Homo sapiens mRNA; cDNA DKFZp586G0321 (from clone DKFZp586G0321)
1609746	-0.4792809 vitelliform macular dystrophy (Best disease, bestrophin)
882248 1917449 2119838	-0.8700731 transgelin -1.0894686 serum amyloid A4, constitutive -0.3045374 a disintegrin-like and metalloprotease (reprolysin type) with thrombospondin type 1 motif, 8
841507	0.0247736 surfactant, pulmonary-associated protein A2
813265	-1.2230435 Homo sapiens mRNA; cDNA DKFZp564H1916 (from clone DKFZp564H1916)
767202	-1.0654145 latent transforming growth factor beta binding protein 2
1156538	0.2252628 potassium inwardly-rectifying channel, subfamily J, member
39600 1630990 41208 322561 951008 841308	-0.6540275 adenylate kinase 5 -0.7312981 ribosomal protein L29 -0.865227 bone morphogenetic protein 1 -0.7970805 ribosomal protein L31 -0.2221875 ESTs -0.9953716 myosin, light polypeptide kinase
sd-71385	

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119290	-0.8224756 cortic al thymocyte receptor (X. laevis CTX) like
344959 810331 1161775 625399	-0.0998837 gene for serine/threonine protein kinase -0.1389612 quiescin Q6 -0.7380353 villin 1 -0.9066615 hypothetical protein similar to beta-transducin family
1470657 160609 665356	-0.3557985 deiodinase, iodothyronine, type II -0.8405669 ESTs -0.1575968 tumor necrosis factor receptor superfamily, member 11b (osteoprotegerin)
1584540	-0.9754693 Homo sapiens mRNA; cDNA DKFZp586M2022 (from clone DKFZp586M2022)
726779 296123	-0.4459955 calponin 1, basic, smooth muscle -0.2814132 Homo sapiens PRO1851 mRNA, complete cds
190059	-0.634814 guanine nucleotide binding protein (G protein), gamma 7
1471829	-0.7275378 uncharacterized hypothalamus protein HSMNP1
2056139 190753 1968422	-1.0673271 LIM domain protein -0.4423669 ESTs -0.5802058 Homo sapiens mRNA full length insert cDNA clone EUROIMAGE 1968422
725390 814826	-0.4620278 glutathione S-transferase pi -0.8443265 ESTs
878836	-1.037352 secretory granule, neuroendocrine protein 1 (7B2 protein)
27769 188388 1597813 1609625 810981 726703	-0.8298821 ESTs -0.6824191 integrin, alpha 10 -0.5406089 hypothetical protein PP1044 -0.5920312 selectin P ligand -0.1805871 hypothetical protein FLJ20699 -0.070084 Homo sapiens clone 23736 mRNA sequence
781014 898222	-1.0161379 suppression of tumorigenicity 5 0.1795078 Homo sapiens clone 24418 mRNA sequence
1475738 1492144 1492147 sd-71385	-0.6753091 ribosomal protein S25 -0.2356446 butyrophilin, subfamily 3, member A2 -0.636656 ribosomal protein S4, X-linked

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1711456 1871116	-0.4896574 H factor (complement)-like 1 -0.9821709 Homo sapiens mRNA; cDNA DKFZp434C1714 (from clone DKFZp434C1714); partial cds
344720 45099	-1.5196431 glycophorin C (Gerbich blood group) -1.3515907 regucalcin (senescence marker protein-30)
307029 25763 2502722	-0.6500872 ribosomal protein L26 -0.2725443 ankylosis, progressive (mouse) homolog -0.3315871 loss of heterozygosity, 11, chromosomal region 2, gene A
1665444 165837	-0.651994 tumor endothelial marker 1 precursor -0.3443143 translocase of inner mitochondrial membrane 22 (yeast) homolog
502518 897963 120138 78946	-0.4342228 laminin, beta 2 (laminin S) -0.9775616 phosphatidic acid phosphatase type 2A 0.1570107 J domain containing protein 1 -0.648366 Homo sapiens pyruvate dehydrogenase kinase 4 mRNA, 3' untranslated region, partial sequence
990881 859192	-0.8335353 -0.2265317 vesicle-associated membrane protein 1 (synaptobrevin 1)
712023 855586	-0.5302372 AT-binding transcription factor 1 -0.3944697 nuclear receptor subfamily 3, group C, member 1
52419 432072	-1.0368509 Friedreich ataxia region gene X123 -0.4193898 nuclear factor of activated T-cells, cytoplasmic, calcineurin-dependent 1
1897947	-0.163146 surfactant, pulmonary-associated protein A2
727229	-0.2668972 mitogen-activated protein kinase kinase kinase 4
810358	-0.5403705 acyl-Coenzyme A dehydrogenase, very long chain
154600 755855	 -0.0683999 phospholipase C, delta 1 -0.6298333 solute carrier family 25 (mitochondrial carrier; adenine nucleotide translocator), member 6
323780	-0.8879854 Homo sapiens cDNA FLJ11177 fis, clone PLACE1007402
230560 283173	-0.1803944 ESTs -0.4293696 EBP50-PDZ interactor of 64 kD
1 71007	

265853	-0.3519023 Homo sapiens cDNA FLJ11298 fis, clone PLACE1009794
161456 184022	-0.8215502 serum amyloid A1 -0.7070266 amyloid beta (A4) precursor protein-binding, family B, member 1 (Fe65)
1636523	-0.1496249 glutathione S-transferase subunit 13 homolog
795730	-0.2892776 signal transduction protein (SH3 containing)
1631682	-0.5162406 peptidylprolyl isomerase E (cyclophilin E) -0.848137
811837 854763	-0.0493902 Homo sapiens cDNA FLJ11341 fis, clone PLACE1010786
50562 46843 178137 1662279	-0.0943363 chromosome 8 open reading frame 4 -0.1736687 neuronal Shc adaptor homolog -0.6839022 ribosomal protein L34 -0.2290724 plakophilin 1 (ectodermal dysplasia/skin fragility syndrome)
1635062 293916 415613 80344 1602798 1910516 740620 277627	-0.8243024 DKFZP586A011 protein -0.3671832 FKBP-associated protein -0.5528761 DHHC1 protein -0.8243418 interleukin 7 receptor -0.4300808 choline kinase-like -0.2845842 ESTs -0.96496 tropomyosin 2 (beta) -0.5329107 Human SH3 domain-containing protein SH3P18 mRNA, complete cds 0.008949 siah binding protein 1; FBP interacting repressor; pyrimidine tract binding splicing factor; Ro ribonucleoprotein-binding
45578 950710	protein 1 -0.1295485 mitogen-activated protein kinase kinase 6 -0.346361 propionyl Coenzyme A carboxylase, alpha polypeptide
768043 308539	-0.4005598 ECSIT -0.6726187 Homo sapiens cDNA FLJ12777 fis, clone NT2RP2001720
595637	-0.5373697 Homo sapiens mRNA; cDNA DKFZp586N012 (from clone DKFZp586N012)
143661 248631	-0.4713778 netrin 4 -0.9106505 aminomethyltransferase (glycine cleavage system protein T)

826622 51981 811848 1533710	0.1339287 KIAA0430 gene product -0.5554908 ribosomal protein L7a -1.1350073 hypothetical protein -0.5988282 ortholog of mouse integral membrane glycoprotein LIG-1
2017756 1637296 50586 877835 773319	-0.2549438 homolog of yeast MOG1 -0.5105214 ribosomal protein S24 -0.6794282 KIAA1545 protein -0.5382125 ribosomal protein L35 -0.285447 ribosomal protein S6 kinase, 70kD, polypeptide 1
2014888 767495 774078 868400 240620	-0.2871529 sushi-repeat protein -0.0366213 GLI-Kruppel family member GLI3 (Greig cephalopolysyndactyly syndrome) -0.6508021 leiomodin 1 (smooth muscle) -0.4928479 glutaminyl-tRNA synthetase -0.5065007 vascular Rab-GAP/TBC-containing
1588791 472186 725143 714472 1854648 855029 197727	-0.1606643 O-6-methylguanine-DNA methyltransferase -0.2687871 RAB32, member RAS oncogene family -0.3410957 hypothetical protein FLJ22418 -0.3677387 KIAA0397 gene product 0.0089186 hemopexin 0.076972 Ac-like transposable element -0.3906866 phosphatidylethanolamine N-methyltransferase
1623016 813841 2783721	-0.6263561 EST 0.0885625 plasminogen activator, tissue -0.6854649 cholinergic receptor, nicotinic, beta polypeptide 1 (muscle)
549933 280907 796181 415233 811920 415415	-0.3547881 interleukin 8 -0.7307519 Kruppel-type zinc finger protein -0.8888496 growth arrest-specific 6 -0.6506593 ribosomal protein L37a -1.2690713 interleukin 11 receptor, alpha -0.646492 EST, Moderately similar to RL1X_HUMAN 60S RIBOSOMAL PROTEIN L18A [H.sapiens]
2018807 154999 68557 647763	-0.2989936 KIAA0468 gene product -0.3876757 hypothetical protein FLJ21007 0.0637586 fatty acid binding protein 1, liver -0.1436566 ESTs
	51981 811848 1533710 2017756 1637296 50586 877835 773319 2014888 767495 774078 868400 240620 1588791 472186 725143 714472 1854648 855029 197727 1623016 813841 2783721 549933 280907 796181 415233 811920 415415

1568967	-0.2938468 ESTs
210717	-0.4522432 syndecan 2 (heparan sulfate proteoglycan 1, cell surface-
	associated, fibroglycan)
1350439	-0.6740833 KIAA0015 gene product
684582	-0.0089016 tryptophanyl tRNA synthetase 2 (mitochondrial)
122394	-0.049399 guanine nucleotide binding protein (G protein), alpha 15 (Gq class)
213136	-0.1139004 BTG family, member 2
625458	-0.2394796 hypothetical protein MGC3234
811162	-0.3821116 fibromodulin
1926246	-0.701769 ESTs, Moderately similar to T46371 hypothetical protein DKFZp434P1018.1 [H.sapiens]
1554167	-1.0818834 hypothetical protein FLJ14529
811088	-1.0851546 ephrin-B3
72778	-1.1752838 caspase 7, apoptosis-related cysteine protease
220293	-0.5078291 Homo sapiens cDNA: FLJ21800 fis, clone HEP00618
1569187	-1.2971252 heparan sulfate (glucosamine) 3-O-sulfotransferase 4
866866	-0.4801351 Ras association (RalGDS/AF-6) domain family 1
81316	-0.6176265 ESTs
1161564	-1.3891189 desmuslin
898044	-0.4691863 metallocarboxypeptidase CPX-1
293001	-0.4472585 hypothetical protein DKFZp434E2318
1570502	-0.3314609 Homo sapiens cDNA FLJ12936 fis, clone NT2RP2005018
839796	-0.6807912 candidate tumor suppressor p33 ING1 homolog
380883	-0.4438655 Homo sapiens cDNA FLJ10158 fis, clone HEMBA1003463
180561	-0.3250633 glutathione S-transferase M4
869450	-0.7659922 ribosomal protein L11
47043	-0.7729643 tensin
810463	-0.0829995 DKFZP566O084 protein
149539	0.4138942 KIAA1700
200354	-0.0613983 thymidine kinase 2, mitochondrial
771173	0.1757755 mitochondrial ribosomal protein S21
270826	-0.9389651 Homo sapiens cDNA FLJ13329 fis, clone OVARC1001795

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	810017 809838 344168 785967	-0.1076821 plasminogen activator, urokinase receptor 0.0956351 -0.386185 polymerase (DNA directed), lambda -0.6049171 erythrocyte membrane protein band 4.1-like 2
	511831 282404	-0.2912672 hypothetical protein MGC12936 -0.5528532 Homo sapiens mRNA for KIAA1671 protein, partial cds
	730036 298231	-0.079736 Mad4 homolog -0.3550893 gamma-aminobutyric acid (GABA) B receptor, 1
	2106144 743880 268234 280776	-1.067765 regulated in glioma -0.3262292 KIAA0263 gene product -0.1535073 Dmx-like 1 -0.2922639 acyl-Coenzyme A dehydrogenase, C-2 to C-3 short chain
F 1 00 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	212078 731308 774471 811837	-0.1831462 integrin, alpha 1 0.5371204 citrate synthase -0.1753902 laminin, beta 1 -0.8764231 eukaryotic translation elongation factor 1 alpha 1
	714437	0.0258281 sema domain, immunoglobulin domain (lg), transmembrane domain (TM) and short cytoplasmic domain, (semaphorin) 4B
	416959 681992	-0.957248 nuclear factor I/B -0.7691299 Homo sapiens cDNA FLJ13384 fis, clone PLACE1001062, highly similar to Homo sapiens mRNA for lysine-ketoglutarate reductase/saccharopine dehydrogenase
	130201 363144	-1.0401997 intercellular adhesion molecule 2 0.1937891 transcription factor AP-2 beta (activating enhancer-binding protein 2 beta)
	857874	-0.3703645 transforming growth factor beta-activated kinase-binding protein 1
	377461 156363 589115	-1.2798035 caveolin 1, caveolae protein, 22kD 0.2685238 hypothetical protein FLJ12934 -0.0115547 matrix metalloproteinase 1 (interstitial collagenase)
	241489 586725	-0.6212938 adrenergic, beta-2-, receptor, surface 0.0683397 protein phosphatase 2, regulatory subunit B (B56), beta isoform

	283124	-0.7933772 ESTs, Moderately similar to LONG-CHAIN FATTY ACID TRANSPORT PROTEIN [M.musculus]
	307328	-0.4022704 hypothetical protein FLJ10948
	37449	-0.3539978 GAS2-related on chromosome 22
	50892	-0.6887497 Homo sapiens, clone MGC:9913 IMAGE:3870821, mRNA, complete cds
	1897944	-0.4855156 ESTs, Weakly similar to T00366 hypothetical protein KIAA0669 [H.sapiens]
E de	377314	-0.561312 casein kinase 2, alpha prime polypeptide
	220851	-0.0351418 crystallin, alpha A
	322233	-0.3025167
	586803	-0.7423407 placental growth factor, vascular endothelial growth factor-related protein
en aprovincia pri ul to ul to un to un approvincia minimalarita	343695	-0.7706809 hypothetical protein FLJ10875
	414999	-0.5002542 ets variant gene 4 (E1A enhancer-binding protein, E1AF)
dive game of the second of the		olocozo iz oto variant gono i (z in tomianosi binanig protom, z in ti
E a	813823	-0.4463267 lumican
	178825	-0.1525189 neurogranin (protein kinase C substrate, RC3)
	85582	-0.0716296 target of myb1 (chicken) homolog-like 2
A second	753620	-0.5248489 insulin-like growth factor binding protein 6
Bari E F	1899338	-0.066168 mannosidase, alpha, class 1A, member 2
nel q p period 5 v od	359250	0.0674486 carbonic anhydrase IV
	345034	-1.0600859 small inducible cytokine subfamily B (Cys-X-Cys), member 14 (BRAK)
	2164744	-0.2307289 neural cell adhesion molecule 1
	613056	-0.3957342 reticulocalbin 1, EF-hand calcium binding domain
	1410444	-0.0368027 amphiregulin (schwannoma-derived growth factor)
	593023	-0.7516846 dystrobrevin, beta
	788511	0.1403148 ribosomal protein S6 kinase, 90kD, polypeptide 1
	455000	
	455269	0.0223607
	1650927	-0.1712163 hypothetical protein DKFZp547E052
	782427	-0.2818886 inhibin, beta B (activin AB beta polypeptide)
	71727	-0.2796517 T-cell acute lymphocytic leukemia 1
	840942	-0.2730317 1-ccil acute lymphocytic leukernia 1 -0.6383764 major histocompatibility complex, class II, DP beta 1
	U7U372	-0.000070- major molocompanionity complex, class II, DF beta 1
	825470	0.5218731 topoisomerase (DNA) II alpha (170kD)

	595213 796694	•	oothetical protein culoviral IAP repeat-containing 5 (survivin)
	810711	0.9128832 ste	aroyl-CoA desaturase (delta-9-desaturase)
	629944 66406 785840	7 1	osin VB bothetical protein DKFZp762E1312 C24 (S. cerevisiae) related gene family, member D
	210862 84295 429182 705064	0.3989864 inte 0.2952867 dol cat	/I-Coenzyme A oxidase 1, palmitoyl erleukin 1 receptor antagonist ichyl-phosphate mannosyltransferase polypeptide 1, alytic subunit nsforming, acidic coiled-coil containing protein 3
	782513	0.4759571 inte	erferon, alpha-inducible protein (clone IFI-6-16)
	839682		iquitin-conjugating enzyme E2N (homologous to yeast
ment and many and many and	768377 2309073 789376 196992	0.3809674 act 0.5226599 friz 0.4339171 thic -0.1015205 ald del	tivity-dependent neuroprotective protein zled (Drosophila) homolog 5 oredoxin reductase 1 lo-keto reductase family 1, member C1 (dihydrodiol hydrogenase 1; 20-alpha (3-alpha)-hydroxysteroid
	785707 1505038 869375	0.4708376 pro 1.1904802 hyp	hydrogenase) otein regulator of cytokinesis 1 pothetical protein FLJ20171 ocitrate dehydrogenase 2 (NADP+), mitochondrial
	109221 345787		AA0286 protein phly expressed in cancer, rich in leucine heptad repeats
	46248		P-ribosyltransferase (NAD+; poly (ADP-ribose) lymerase)
	79520 469383 509588	0.6769129 RA 0.5558732 chr 0.4157059 TA	AB2, member RAS oncogene family romosome 8 open reading frame 1 ATA box binding protein (TBP)-associated factor, RNA lymerase II, J, 20kD
	292936 686172 788655	•	pothetical protein FLJ10468 AA0008 gene product PAP protein

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770992 1587847	0.3331315 0.1732392 minichromosome maintenance deficient (mis5, S. pombe) 6
79710 2056566	0.4579008 KIAA0174 gene product 0.2341905 integrin, alpha M (complement component receptor 3, alpha; also known as CD11b (p170), macrophage antigen alpha polypeptide)
454339 594226	0.3738929 thiopurine S-methyltransferase 0.403676 Homo sapiens cDNA FLJ14459 fis, clone HEMBB1002409
897813	0.4749348 polyadenylate binding protein-interacting protein 1
824694	0.0268936 protein tyrosine phosphatase type IVA, member 1
725454 79254 1472719	0.6669973 CDC28 protein kinase 2 0.0732432 MHC class I region ORF 0.2550065 SMT3 (suppressor of mif two 3, yeast) homolog 1
2054635	0.7914755 proteasome (prosome, macropain) subunit, alpha type, 7
289978 155806	0.6308861 ubiquitin-like 4 0.2747101 2'-5'-oligoadenylate synthetase 2 (69-71 kD)
147834 26171 2322367 769921 73009	0.4412795 zinc finger protein 217 0.3656147 KIAA0856 protein 0.5832711 reticulon 4 0.6464245 ubiquitin carrier protein E2-C 0.4822587 ESTs, Weakly similar to A43932 mucin 2 precursor, intestinal
2014034	[H.sapiens] 0.6620303 methylene tetrahydrofolate dehydrogenase (NAD+ dependent), methenyltetrahydrofolate cyclohydrolase
359887	0.8692353 translocase of inner mitochondrial membrane 17 (yeast) homolog A
488964 1476065 1422338 268946 686552 149544 770675	0.9569176 H2A histone family, member O 0.289904 stathmin 1/oncoprotein 18 0.5874097 ribonucleotide reductase M2 polypeptide 0.5440003 WD40 protein Ciao1 0.7231319 golgi phosphoprotein 1 0.283642 neuroepithelial cell transforming gene 1 0.454729 Homo sapiens cDNA: FLJ21323 fis, clone COL02374
	79710 2056566 454339 594226 897813 824694 725454 79254 1472719 2054635 289978 155806 147834 26171 2322367 769921 73009 2014034 359887 488964 1476065 1422338 268946 686552 149544

346257	0.1515899 minichromosome maintenance deficient (S. cerevisiae) 4
429799	0.6915155 hypothetical protein FLJ21939 similar to 5-azacytidine induced gene 2
143997	0.1883447 proteasome (prosome, macropain) 26S subunit, non- ATPase, 10
122241	0.7394284 proteasome (prosome, macropain) subunit, beta type, 2
823598	0.9153521 proteasome (prosome, macropain) 26S subunit, non- ATPase, 12
814632	0.0960611 splicing factor, arginine/serine-rich (transformer 2 Drosophila homolog) 10
810316	0.3732635 very long-chain acyl-CoA synthetase; lipidosin
290841	0.8346933 H2B histone family, member A
347373	0.7563599 transcription elongation factor B (SIII), polypeptide 1 (15kD, elongin C)
700792	0.4949149 cyclin-dependent kinase inhibitor 3 (CDK2-associated dual specificity phosphatase)
1554549	0.9764206 hydroxyacyl glutathione hydrolase
897770	0.4270685
504308	
132828	0.2875519 Down syndrome critical region gene 1-like 2
292388	0.112577
1616253	0.567837 breast carcinoma amplified sequence 1
796469	1.0269115 HSPC150 protein similar to ubiquitin-conjugating enzyme
2139152	-0.4373776 Homo sapiens clone 24473 mRNA sequence
130276	0.2154295 Homo sapiens mRNA; cDNA DKFZp586H0324 (from clone DKFZp586H0324)
34149	0.309569 KIAA0227 protein
150003	
287749	
25380	0.164077 ESTs, Moderately similar to JC5238 galactosylceramide-like protein, GCP [H.sapiens]
745083	0.4107735 ubiquitin specific protease 18
810899	
768059	
sd-71385	•
00	

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815026	0.4472842 cleavage stimulation factor, 3' pre-RNA, subunit 3, 77kD
741474 788641	0.7732047 glucose phosphate isomerase 0.3638059 adaptor-related protein complex 1, sigma 2 subunit
2043167 1492780	0.7167431 BCL2-associated athanogene 3 0.6544659 Homo sapiens cDNA FLJ14459 fis, clone HEMBB1002409
2306987 754653	0.4402912 secreted and transmembrane 1 0.4405873 cadherin, EGF LAG seven-pass G-type receptor 3, flamingo
743810 126858	(Drosophila) homolog 0.3667087 hypothetical protein MGC2577 0.2389814 sterol O-acyltransferase (acyl-Coenzyme A: cholesterol
2017415	acyltransferase) 1 0.43546 centromere protein A (17kD)
280507	0.2846518 hypoxanthine phosphoribosyltransferase 1 (Lesch-Nyhan syndrome)
67237 43833 42076	0.1406869 ESTs -0.1197546 diacylglycerol kinase, gamma (90kD) 0.3194462 TRK-fused gene
531319 784129	0.1093932 serine/threonine kinase 12 0.2902598 tyrosine 3-monooxygenase/tryptophan 5-monooxygenase
586895	activation protein, gamma polypeptide 0.2549108 small nuclear ribonucleoprotein polypeptide G

Example VIII: Genes for discriminating between DCIS Grade I and Grade III

As shown in Table 7 below, 350 genes were identified as being able to discriminate between grades of DCIS. The actual data corresponding to this table is shown in Figure 4.

Table 7

CloneID	Weight	Description
2460159	4.6268975 tyrosine kinase, non-re	eceptor, 1
358151	2.8783989 zinc finger protein 33a	(KOX 31)

795382	2.772572 Rap1 guanine-nucleotide-exchange factor directly activated by cAMP
714472	2.6829714 KIAA0397 gene product
725649	2.6433625 nuclear factor of activated T-cells, cytoplasmic, calcineurin- dependent 4
51218	2.6200167 ESTs
504959	2.5777963 Homo sapiens mRNA; cDNA DKFZp586G0321 (from clone DKFZp586G0321)
647397	2.5320153 ESTs
_	2.4198892 plakophilin 4
279720	2.4136316 ESTs, Moderately similar to A47582 B-cell growth factor precursor [H.sapiens]
298231	2.4096172 gamma-aminobutyric acid (GABA) B receptor, 1
172783	2.3339699 hypothetical protein FLJ10390
261609	2.311051 ESTs, Weakly similar to S65657 alpha-1C-adrenergic receptor splice form 2 [H.sapiens]
826668	2.2951137 KIAA0274 gene product
1493383	2.2616737 Homo sapiens mRNA; cDNA DKFZp434H2418 (from clone
	DKFZp434H2418)
0047750	0.0400000 L
2017756	2.2403622 homolog of yeast MOG1
1455566	2.1756946 adenosine A3 receptor
725321	2.1658095 estrogen receptor 1
180561	2.1393944 glutathione S-transferase M4 2.1165022 Homo sapiens mRNA; cDNA DKFZp586P1124 (from clone
32050	DKFZp586P1124)
215000	2.1150123 vasoactive intestinal peptide receptor 1
2019750	2.0957741 SEC14 (S. cerevisiae)-like 2
283124	2.0921779 ESTs, Moderately similar to LONG-CHAIN FATTY ACID
	TRANSPORT PROTEIN [M.musculus]
490615	2.0635424 tubulin, gamma 2
666138	2.0625667 hypothetical protein DKFZp761J1523
418129	2.0541212 nuclear mitotic apparatus protein 1
1733262	2.052666 BLu protein
1588791	2.044069 O-6-methylguanine-DNA methyltransferase
101-01	O COMONO DA LA CAMBILLA DA CAMBILLA DE COMONA
461761	2.035953 angiogenin, ribonuclease, RNase A family, 5

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	1.9930012 kinesin protein 9 gene 1.9781057 Homo sapiens mRNA; cDNA DKFZp566H0124 (from clone DKFZp566H0124)
28643 470261 1630990 810981 767495	 1.9733043 hypothetical protein DKFZp564D1378 1.9160486 1.9155421 ribosomal protein L29 1.9132205 hypothetical protein FLJ20699 1.9052671 GLI-Kruppel family member GLI3 (Greig cephalopolysyndactyly syndrome) 1.8996792 tumor necrosis factor (ligand) superfamily, member 13
1572196	1.8855996 secreted modular calcium-binding protein 2
1706635	1.8754346 bone gamma-carboxyglutamate (gla) protein (osteocalcin)
186301	1.8424458 Homo sapiens cDNA FLJ12924 fis, clone NT2RP2004709
726703	1.842399 Homo sapiens clone 23736 mRNA sequence
214205	1.8416657 Homo sapiens, clone MGC:17687 IMAGE:3865868, mRNA, complete cds
784178	1.822159 Homo sapiens mRNA; cDNA DKFZp586M0723 (from clone DKFZp586M0723)
346902	1.8196723 ESTs, Weakly similar to ALU2_HUMAN ALU SUBFAMILY SB SEQUENCE CONTAMINATION WARNING ENTRY [H.sapiens]
1367678 190059	1.8184255 KIAA0356 gene product 1.7988757 guanine nucleotide binding protein (G protein), gamma 7
1456937	1.7980645 oviductal glycoprotein 1, 120kD (mucin 9, oviductin)
45578 248631	1.7935215 mitogen-activated protein kinase kinase 6 1.7701919 aminomethyltransferase (glycine cleavage system protein T)
1562231 154466	1.7668811 SET binding protein 1 1.7516237 STIP1 homology and U-Box containing protein 1
2524445 277266	1.7392197 neuronal PAS domain protein 1 1.7307752 Homo sapiens, clone IMAGE:3625550, mRNA, partial cds
741891 sd-71385	1.7297992 RAB2, member RAS oncogene family-like

206217	1.7227658 nuclear receptor subfamily 1, group H, member 3
2028876 730036 1558233 502518 356835 744994 810358	 1.7128178 splicing factor, arginine/serine-rich 5 1.70573 Mad4 homolog 1.7035129 ESTs 1.7027287 laminin, beta 2 (laminin S) 1.7014349 hypothetical protein MGC10500 1.6938348 hypothetical protein FLJ12242 1.6924263 acyl-Coenzyme A dehydrogenase, very long chain
813854 768043 264632 505243 344073	 1.6914133 purine-rich element binding protein A 1.6854123 ECSIT 1.6591646 ESTs 1.6581518 inositol 1,4,5-triphosphate receptor, type 2 1.6571604 ESTs, Weakly similar to K1CI_HUMAN KERATIN, TYPE I CYTOSKELETAL 9 [H.sapiens]
343760	1.6435915 SH3 domain binding glutamic acid-rich protein like 2
1569418	1.6408767 Homo sapiens cDNA FLJ11385 fis, clone HEMBA1000520
202577 823634 839796	1.6393518 histamine N-methyltransferase 1.6366047 ESTs 1.630887 candidate tumor suppressor p33 ING1 homolog
183440 344959 1574252 669359	1.6297915 arylsulfatase A 1.6268043 gene for serine/threonine protein kinase 1.6211224 DKFZP586D0623 protein 1.6174855 Homo sapiens clone 24405 mRNA sequence
74070 1660649 796723	1.6096611 endosulfine alpha 1.6051582 suppressor of white apricot homolog 2 1.6041219 Homo sapiens clone CDABP0014 mRNA sequence
743146 789147 342181 33076 782497	1.597094 hypothetical protein FLJ23403 1.5937127 enolase 2, (gamma, neuronal) 1.5926355 B-cell CLL/lymphoma 2 1.5824352 cholinephosphotransferase 1 1.5804781 Homo sapiens, clone IMAGE:3010666, mRNA, partial cds
470261 809507 712460 sd-71385	1.5751564 SMA5 1.5723669 hypothetical protein FLJ20568 1.5721924 natural killer-tumor recognition sequence

293569 1518402 155072 455269 262804 1492238 364865 2325804	1.5627158 KIAA1361 protein 1.5557559 ESTs 1.5538593 1.5532348 hypothetical protein MGC2941 1.5484044 HSPC003 protein 1.5483958 hypothetical protein FLJ21062
1635062 344168 1517171 769600 325583 814826 1570502	1.5421405 polymerase (DNA directed), lambda 1.5381921 interleukin 2 receptor, alpha 1.5335878 uracil-DNA glycosylase 2 1.5245817 EST 1.5223296 ESTs
511831 124922 2072768 2021882 627248 725503 285312 141731 1456701 898222	1.5183298 hypothetical protein MGC12936 1.5052004 KRAB-zinc finger protein SZF1-1 1.4886791 nuclear receptor coactivator 3 1.4854251 sodium channel, nonvoltage-gated 1, delta 1.4827018 SBBI31 protein 1.476461 D-dopachrome tautomerase 1.4699425 1.469088 1.4668054 B-cell CLL/lymphoma 9 1.4667947 Homo sapiens clone 24418 mRNA sequence
725284 154999 1592530 590310	 1.4638006 phosphorylase kinase, gamma 2 (testis) 1.4564204 hypothetical protein FLJ21007 1.4558873 mammalian inositol hexakisphosphate kinase 2 1.4534138 Homo sapiens mRNA; cDNA DKFZp434E2321 (from clone DKFZp434E2321); partial cds
416042	1.446691 ESTs, Weakly similar to ALU1_HUMAN ALU SUBFAMILY J SEQUENCE CONTAMINATION WARNING ENTRY [H.sapiens]
2017144 309449 450301	1.4450059 CGI-41 protein 1.4412699 ribosomal protein S4, Y-linked 1.4403678 mutL (E. coli) homolog 3
cd_71385	

1492468 121454 743880 1568967 1568989 25274 2413337	1.4375076 DEME-6 protein 1.4361788 arachidonate 12-lipoxygenase 1.4294838 KIAA0263 gene product 1.4109448 ESTs 1.4094472 ESTs 1.4078252 g20 protein 1.4058288 sortilin-related receptor, L(DLR class) A repeats-containing
197913	1.3994468 splicing factor proline/glutamine rich (polypyrimidine tract- binding protein-associated)
292770	1.3952057 Homo sapiens, clone IMAGE:3627860, mRNA, partial cds
143332 809779 138242	1.3875805 neuropeptide Y receptor Y1 1.3848708 KIAA0239 protein 1.3820433 ESTs, Moderately similar to MAS2_HUMAN MANNAN-BINDING LECTIN SERINE PROTEASE 2 PRECURSOR [H.sapiens]
270127 49240 826622 1858837 1583198 345858 208387 502782 26294 669379	 1.3734143 KIAA0430 gene product 1.3719341 ESTs 1.3712112 ESTs, Weakly similar to S65824 reverse transcriptase homolog [H.sapiens] 1.3629187 cisplatin resistance associated 1.3597205 KIAA1407 protein 1.3589088 RAN binding protein 3
61061 49630	1.3450626 fibromodulin 1.3437368 hypothetical protein FLJ20585 1.3412751 calcium channel, voltage-dependent, L type, alpha 1D subunit
825659 769921	-2.8345933 N-myc downstream regulated -2.712812 ubiquitin carrier protein E2-C

- 1476053 -2.529039 RAD51 (S. cerevisiae) homolog (E coli RecA homolog)
- 809557 -2.4910488 minichromosome maintenance deficient (S. cerevisiae) 3
- 150897 -2.4302462 UDP-GlcNAc:betaGal beta-1,3-N-

acetylglucosaminyltransferase 3

- 814526 -2.37273 seb4D
- 796694 -2.3104815 baculoviral IAP repeat-containing 5 (survivin)
- 814792 -2.3001334 ubiquitin specific protease 10
- 1536236 -2.2971152 hypothetical protein FLJ13154
- 2017415 -2.2748559 centromere protein A (17kD)
- 210862 -2.2408114 acyl-Coenzyme A oxidase 1, palmitoyl
- 795543 -2.2403389 thioredoxin peroxidase (antioxidant enzyme)
- 773301 -2.2239527 cadherin 3, type 1, P-cadherin (placental)
- 700792 -2.2203557 cyclin-dependent kinase inhibitor 3 (CDK2-associated dual specificity phosphatase)
- 1883327 -2.2093248 ESTs

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- 1523225 -2.1936598 oncostatin M receptor
- 208718 -2.1882654 annexin A1
- 1702742 -2.1567809 solute carrier family 7 (cationic amino acid transporter, y+ system), member 5
- 753378 -2.151145 hypothetical protein FLJ22649 similar to signal peptidase SPC22/23
- 429222 -2.1351022 CGI-107 protein
- 292388 -2.1343367
- 781047 -2.1262682 budding uninhibited by benzimidazoles 1 (yeast homolog)
- 450854 -2.0819596 craniofacial development protein 1
- 823598 -2.0693353 proteasome (prosome, macropain) 26S subunit, non-ATPase,
- 1422338 -2.0689327 ribonucleotide reductase M2 polypeptide
- 705064 -2.0601901 transforming, acidic coiled-coil containing protein 3
- 770675 -2.0272207 Homo sapiens cDNA: FLJ21323 fis, clone COL02374
- 345787 -1.9992897 highly expressed in cancer, rich in leucine heptad repeats
- 471196 -1.9985751 integral membrane protein 3

sd-71385

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753215 -1.9954476 guanine nucleotide binding protein (G protein), alpha inhibiting
                    activity polypeptide 1
869375 -1.9803724 isocitrate dehydrogenase 2 (NADP+), mitochondrial
842818 -1.9796507 lysyl-tRNA synthetase
229579 -1.9782493 Golgi apparatus protein 1
1916461 -1.9736332 hypothetical protein
 129294 -1.9730334 ESTs
789376 -1.9719575 thioredoxin reductase 1
1897302 -1.9681938 serine/threonine kinase 17b (apoptosis-inducing)
951117 -1.9577807 eukaryotic translation elongation factor 1 gamma
1642496 -1.9484342 hypothetical protein MGC11266
 591465 -1.9277062 Homo sapiens, clone MGC:2908 IMAGE:3029644, mRNA,
                    complete cds
 149355 -1.8999907 translocating chain-associating membrane protein
 259950 -1.8958779 hypothetical protein FLJ14991
 897770 -1.8887034
 878798 -1.8774101 beta-2-microglobulin
 746229 -1.873434 mitogen-activated protein kinase kinase kinase kinase 4
 624867 -1.8594915 hypothetical protein FLJ20186
 504308 -1.8590049 hypothetical protein FLJ10540
 727251 -1.8553757 CD9 antigen (p24)
 897774 -1.8532125 adenine phosphoribosyltransferase
1901310 -1.8434048 KIAA1209 protein
 292936 -1.838542 hypothetical protein FLJ10468
1518591 -1.8382765
 321354 -1.8123042 mitochondrial ribosomal protein L15
1903066 -1.8071117 keratin, hair, basic, 1
 235180 -1.8004039 ESTs, Moderately similar to ALU1_HUMAN ALU SUBFAMILY
                    J SEQUENCE CONTAMINATION WARNING ENTRY
                    [H.sapiens]
 740604 -1.7995578 interferon stimulated gene (20kD)
 233464 -1.7978669 epithelial V-like antigen 1
 951241 -1.7961406 clone HQ0310 PRO0310p1
1587847 -1.7886092 minichromosome maintenance deficient (mis5, S. pombe) 6
 624390 -1.7800152 DC13 protein
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128711 -1.7791673 anillin (Drosophila Scraps homolog), actin binding protein
 308633 -1.7667341 hypothetical protein FLJ10339
 884425 -1.761486 chaperonin containing TCP1, subunit 5 (epsilon)
 745394 -1.7583344 Homo sapiens cDNA: FLJ23249 fis, clone COL04196
 852829 -1.7579704 karyopherin alpha 3 (importin alpha 4)
 122241 -1.7490794 proteasome (prosome, macropain) subunit, beta type, 2
 307255 -1.7294781 basement membrane-induced gene
  32493 -1.7273675 integrin, alpha 6
 454896 -1.7216328 DnaJ (Hsp40) homolog, subfamily A, member 2
1876217 -1.7186822 DnaJ (Hsp40) homolog, subfamily A, member 2
 815556 -1.7161709 hypothetical protein FLJ10430
 589869 -1.7120206 transcriptional co-activator with PDZ-binding motif (TAZ)
2012523 -1.7104314 fatty acid binding protein 5 (psoriasis-associated)
1946448 -1.7092537 caveolin 2
 841370 -1.7047297 glutamic-oxaloacetic transaminase 2, mitochondrial (aspartate
                    aminotransferase 2)
1600239 -1.6989076 HSPC037 protein
 462926 -1.6981004 NIMA (never in mitosis gene a)-related kinase 2
 144880 -1.6980733 hypothetical protein from EUROIMAGE 1759349
 744047 -1.6910803 polo (Drosophia)-like kinase
 624627 -1.6888444 ribonucleotide reductase M2 polypeptide
 788566 -1.6752673 Purkinje cell protein 4
 858293 -1.6732706 tyrosine 3-monooxygenase/tryptophan 5-monooxygenase
                    activation protein, theta polypeptide
 470148 -1.659943 ESTs
  66902 -1.6546363 ESTs
  77533 -1.6505418 inositol polyphosphate-5-phosphatase, 40kD
 825282 -1.6417354 DKFZP586L0724 protein
 785840 -1.6406565 SEC24 (S. cerevisiae) related gene family, member D
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824962	-1.6399941 karyopherin alpha 2 (RAG cohort 1, importin alpha 1)
	-1.6396499 eyes absent (Drosophila) homolog 2 -1.6264151 minichromosome maintenance deficient (S. cerevisiae) 2 (mitotin)
2054635	-1.6254343 proteasome (prosome, macropain) subunit, alpha type, 7
267816	-1.6215197 syndecan binding protein (syntenin) -1.6173276 KIAA0551 protein -1.6083862 origin recognition complex, subunit 6 (yeast homolog)-like
796469	-1.6065509 HSPC150 protein similar to ubiquitin-conjugating enzyme
813256	-1.6027042 ATP-binding cassette, sub-family B (MDR/TAP), member 1
811590 823756	-1.60128 chloride intracellular channel 1 -1.6006729 protease, serine, 2 (trypsin 2) -1.6001102 hypothetical protein FLJ11100 -1.5995448 mitogen inducible 2 -1.597146 capping protein (actin filament) muscle Z-line, alpha 2
377368 1506046	-1.5960999 hypothetical protein MGC2577 -1.5955102 cell death regulator aven -1.5918448 hypothetical protein FLJ10815 -1.5890389 polymyositis/scleroderma autoantigen 1 (75kD)
1604703	-1.5853167 major histocompatibility complex, class I, F
429182	-1.5765159 GATA-binding protein 6 -1.5699689 dolichyl-phosphate mannosyltransferase polypeptide 1, catalytic subunit
826355	-1.5677013 vesicle-associated membrane protein 5 (myobrevin)
746190 131091 665384	-1.5659923 KIAA0227 protein -1.5635857 hypothetical protein DKFZp761B1514 -1.558448 Not56 (D. melanogaster)-like protein -1.5550415 KIAA1609 protein -1.5501423 neighbor of COX4 -1.544427 uncharacterized bone marrow protein BM040
949988	-1.5437326 Homo sapiens mRNA; cDNA DKFZp586E1124 (from clone DKFZp586E1124); complete cds

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133213	-1.5433392 fucosyltransferase 4 (alpha (1,3) fucosyltransferase, myeloid-specific)
1660666	-1.5423907 carbonic anhydrase VB, mitochondrial
	-1.5372729 ESTs
1614140	-1.5341302 Ris
201890	-1.5333626 baculoviral IAP repeat-containing 3
38925	-1.5304604 ESTs, Moderately similar to A47582 B-cell growth factor precursor [H.sapiens]
511850	-1.5294752 proteasome (prosome, macropain) 26S subunit, ATPase, 1
489489	-1.5169076 lamin B receptor
825470	-1.5154306 topoisomerase (DNA) II alpha (170kD)
42831	-1.5136459 N-terminal kinase-like
	-1.5066053 kallikrein 6 (neurosin, zyme)
730410	-1.506168 lymphocyte-specific protein tyrosine kinase
810983	-1.5060386 DKFZP434H132 protein
731223	-1.5058799 proteasome (prosome, macropain) subunit, beta type, 2
259017	-1.5053827 ESTs
340745	-1.5041978 ESTs
746163	-1.5000358 ESTs, Weakly similar to ALU1_HUMAN ALU SUBFAMILY J SEQUENCE CONTAMINATION WARNING ENTRY [H.sapiens]
1466621	-1.4998321 ATPase, Ca++ transporting, type 2C, member 1
809588	-1.49754 gamma-glutamyl hydrolase (conjugase, folylpolygammaglutamyl hydrolase)
813707	-1.4946872 regulator of G-protein signalling 16
188335	-1.4895328 egf-like module containing, mucin-like, hormone receptor-like sequence 2
1493160	-1.4851884 small inducible cytokine subfamily B (Cys-X-Cys), member 10
531886	-1.4842053 Homo sapiens clone FLB3344 PRO0845 mRNA, complete cds
753428	-1.4835152 Homo sapiens, clone IMAGE:3542597, mRNA, partial cds
897731	-1.4833877 latrophilin
	-1.4811859 ETAA16 protein
	-1.4755682 glutathione S-transferase A3
	-1.4735772 ADP-ribosylation factor-like 7

785368 -1.4680554 PDZ-binding kinase; T-cell originated protein kinase 825606 -1.4667088 kinesin-like 1 531319 -1.4659871 serine/threonine kinase 12 66406 -1.4641179 hypothetical protein DKFZp762E1312 470124 -1.4635711 RAD1 (S. pombe) homolog

Example IX: Genes for discriminating between IDC Grade I and Grade III

As shown in Table 8 below, 300 genes were identified as being able to discriminate between two grades of IDC.

Table 8

CloneID 1706635	Weight Description 3.279305 bone gamma-carboxyglutamate (gla) protein (osteocalcin)
666138 795382	3.1366533 hypothetical protein DKFZp761J1523 3.0005724 Rap1 guanine-nucleotide-exchange factor directly activated by cAMP
1500542 270127	2.8512705 hypothetical protein FLJ21062 2.8021264 regulator of G-protein signalling 11 2.7455118
588262	2.7127609 Homo sapiens, Similar to RIKEN cDNA 2600001A11 gene, clone MGC:9907 IMAGE:3870073, mRNA, complete cds
277266	2.5004627 Homo sapiens, clone IMAGE:3625550, mRNA, partial cds
2090129	2.4882736 chromobox homolog 2 (Drosophila Pc class)
325583 358151 786675	2.4558352 EST2.4309122 zinc finger protein 33a (KOX 31)2.3877233 epididymis-specific, whey-acidic protein type, four-disulfide core; putative ovarian carcinoma marker
854763	2.378956 Homo sapiens cDNA FLJ11341 fis, clone PLACE1010786
1592976	2.3417116 microphthalmia-associated transcription factor
sd-71385	

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1562231 767176	2.286418 SET binding protein 1 2.2472282 tumor necrosis factor (ligand) superfamily, member 13
784178	2.2026739 Homo sapiens mRNA; cDNA DKFZp586M0723 (from clone DKFZp586M0723)
811162	2.1926501 fibromodulin
	2.1886432 laminin, beta 2 (laminin S)
754429	KIAA0512 [H.sapiens]
214205	2.1127961 Homo sapiens, clone MGC:17687 IMAGE:3865868, mRNA, complete cds
61061	2.1046094 hypothetical protein FLJ20585
344959	2.0756965 gene for serine/threonine protein kinase
814815	·
262804	·
1455566	2.0489061 adenosine A3 receptor
1469149	1.9778151 Homo sapiens clone 24606 mRNA sequence
1403143	1.97701011101110 Sapieris cione 24000 mixiva sequence
358217	1.9703763 glypican 4
206217	1.9595484 nuclear receptor subfamily 1, group H, member 3
261609	1.958083 ESTs, Weakly similar to S65657 alpha-1C-adrenergic receptor splice form 2 [H.sapiens]
197525	1.9550731 flavin containing monooxygenase 5
857640	1.9532398 collagen, type VI, alpha 2
812143	1.9469226 fibronectin leucine rich transmembrane protein 3
012110	1.0 100220 haromodin rodonio non tranomonara protoni o
898222	1.9409805 Homo sapiens clone 24418 mRNA sequence
418129	1.9385868 nuclear mitotic apparatus protein 1
2021882	1.9213983 sodium channel, nonvoltage-gated 1, delta
293819	1.919948 oxidoreductase UCPA
203003	1.8989675 non-metastatic cells 4, protein expressed in
200000	1.0000070 Hori-inclastatic cells 4, protein expressed in
705274	1.8986548 diacylglycerol kinase, delta (130kD)
124922	1.8975916 KRAB-zinc finger protein SZF1-1
2505310	1.8859615 calcium/calmodulin-dependent protein kinase I
1492238	1.8795818 HSPC003 protein
810358	1.8732889 acyl-Coenzyme A dehydrogenase, very long chain

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2391494	1.8698169 ephrin-A4
1592530	1.8669954 mammalian inositol hexakisphosphate kinase 2
810671 2108048 730036 810741 45578 647397 767495	 1.8659671 hypothetical protein FLJ22269 1.8648249 DNB5 1.8564918 Mad4 homolog 1.831265 GABA(A) receptor-associated protein 1.8274273 mitogen-activated protein kinase kinase 6 1.8263711 ESTs 1.8149096 GLI-Kruppel family member GLI3 (Greig cephalopolysyndactyly syndrome) 1.8117146 Homo sapiens mRNA; cDNA DKFZp586P1124 (from clone DKFZp586P1124)
811848	1.8106436 hypothetical protein
186301	1.8021505 Homo sapiens cDNA FLJ12924 fis, clone NT2RP2004709
366526	1.7836259 ESTs, Weakly similar to ALU5_HUMAN ALU SUBFAMILY SC SEQUENCE CONTAMINATION WARNING ENTRY [H.sapiens]
813154	1.7815662 nuclear factor I/A
85195	1.778695 growth arrest and DNA-damage-inducible, gamma
2460159 504959	1.7730834 tyrosine kinase, non-receptor, 1 1.7723846 Homo sapiens mRNA; cDNA DKFZp586G0321 (from clone DKFZp586G0321)
742094	1.7684813 hypothetical protein FLJ20950
256619	1.7658619 hydroxysteroid (17-beta) dehydrogenase 7
726699	1.7594888 ESTs, Weakly similar to AAB47496 NG5 [H.sapiens]
2017144 74070 1762111	1.7514699 CGI-41 protein1.7336683 endosulfine alpha1.7236006 natriuretic peptide receptor C/guanylate cyclase C (atrionatriuretic peptide receptor C)
795750	1.7116475 Homo sapiens clone 25056 mRNA sequence
2019750	1.7095011 SEC14 (S. cerevisiae)-like 2
2325804	1.7039903 95 kDa retinoblastoma protein binding protein

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279720	1.693594 ESTs, Moderately similar to A47582 B-cell growth factor precursor [H.sapiens]
132857	1.6875688 Homo sapiens mRNA; cDNA DKFZp586N1323 (from clone DKFZp586N1323)
1898619 681992	1.684954 hypothetical protein MGC15737 1.6802347 Homo sapiens cDNA FLJ13384 fis, clone PLACE1001062, highly similar to Homo sapiens mRNA for lysine-ketoglutarate
	reductase/saccharopine dehydrogenase
85450	1.6753438 acyl-Coenzyme A oxidase 2, branched chain
51218	1.6748931 ESTs
1925280	1.6563658 homologous to yeast nitrogen permease (candidate tumor suppressor)
250883	1.6504115 ubiquitin-activating enzyme E1-like
172783	1.6502064 hypothetical protein FLJ10390
208387	1.6498608 KIAA1407 protein
703964	1.6452497 inositol polyphosphate phosphatase-like 1
298231	1.6428171 gamma-aminobutyric acid (GABA) B receptor, 1
277848	1.6407233 ADP-ribosylation factor 6
669359	1.6386236 Homo sapiens clone 24405 mRNA sequence
321455	1.6327309 Homo sapiens, Similar to RIKEN cDNA 1110002C08 gene, clone MGC:9564 IMAGE:3872267, mRNA, complete cds
796152	1.6295159 Homo sapiens cDNA FLJ11685 fis, clone HEMBA1004934
502782	1.614818 RAN binding protein 3
360778	1.6091782
950574	1.606032 H3 histone, family 3B (H3.3B)
788334	1.5816572 mitochondrial ribosomal protein L23
782497	1.5767626 Homo sapiens, clone IMAGE:3010666, mRNA, partial cds
1733262	1.5656438 BLu protein
342181	1.5642436 B-cell CLL/lymphoma 2
1632248	1.5590794 Homo sapiens cDNA FLJ14181 fis, clone NT2RP2004300
1526826	1.5576997 homeo box B2
145132	1.5520317 mannose-P-dolichol utilization defect 1
183062	1.5400324 ubiquitin specific protease 21
	•

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823663	1.5337257 fragile X mental retardation, autosomal homolog 2
190059	1.5258594 guanine nucleotide binding protein (G protein), gamma 7
111721 840882 1902764	1.5045285 insulin induced protein 2 1.5039692 nucleotide binding protein 1.5037016 6-phosphofructo-2-kinase/fructose-2,6-biphosphatase 3
414999	1.5021506 ets variant gene 4 (E1A enhancer-binding protein, E1AF)
1636523	1.4948923 glutathione S-transferase subunit 13 homolog
594683 1916575	1.4862053 ESTs 1.4825298 BCL2-interacting killer (apoptosis-inducing)
1456937	1.4730662 oviductal glycoprotein 1, 120kD (mucin 9, oviductin)
25274 795288	1.464238 g20 protein 1.4625618 ubiquitin specific protease 4 (proto-oncogene)
1569418	1.4599089 Homo sapiens cDNA FLJ11385 fis, clone HEMBA1000520
753700 1691237 741891 1493383	 1.4522717 Ras-related GTP-binding protein 1.4516493 ESTs 1.450795 RAB2, member RAS oncogene family-like 1.4468163 Homo sapiens mRNA; cDNA DKFZp434H2418 (from clone DKFZp434H2418)
1675273 293916 39600 1573087 839796	1.445145 RAR-related orphan receptor C 1.4265341 FKBP-associated protein 1.4224626 adenylate kinase 5 1.4200341 KIAA0592 protein 1.4114662 candidate tumor suppressor p33 ING1 homolog
296123	1.4057088 Homo sapiens PRO1851 mRNA, complete cds
727263	1.4046408 heterogeneous nuclear ribonucleoprotein U (scaffold
265045	attachment factor A) 1.4041143 Homo sapiens mRNA; cDNA DKFZp586B0918 (from clone DKFZp586B0918)
869450 714472	1.4034983 ribosomal protein L11 1.4006155 KIAA0397 gene product

825296	1.3963737 low density lipoprotein receptor defect C complementing
753301	1.3962897 carcinoembryonic antigen-related cell adhesion molecule 1 (biliary glycoprotein)
700527	
811565	•
1630990	·
70749	1.3810753 Homo sapiens cDNA: FLJ21874 fis, clone HEP02488
814826	1.3785178 ESTs
175103	1.376581 cadherin, EGF LAG seven-pass G-type receptor 2, flamingo (Drosophila) homolog
215000	1.3677098 vasoactive intestinal peptide receptor 1
810331	1.3656069 quiescin Q6
1681421	1.3645145 EGF-like-domain, multiple 3
502198	1.3619124 protein phosphatase 1, regulatory (inhibitor) subunit 5
825365	1.361275 hypothetical protein FLJ21919
	1.3582677 thrombospondin 3
2067500	·
490615	·
743880	1.3528093 KIAA0263 gene product
812099	1.3444147 RNA binding motif protein 5
868652	1.3432331 complement component 4B
120138	1.3384804 J domain containing protein 1
156363	1.3358314 hypothetical protein FLJ12934
813584	, , , , , , , , , , , , , , , , , , ,
726703	1.3247883 Homo sapiens clone 23736 mRNA sequence
809507	1.3225829 hypothetical protein FLJ20568
2018423	1.3212314 death-associated protein kinase 2
292806	1.3201283 chromosome segregation 1 (yeast homolog)-like
741977	1.3200635 B-factor, properdin
610326	-4.5776738
	-3.6341126 core-binding factor, beta subunit
	-3.6328144 karyopherin alpha 2 (RAG cohort 1, importin alpha 1)
825659	-3.6247426 N-myc downstream regulated
796694	-3.5038212 baculoviral IAP repeat-containing 5 (survivin)

149355	-3.4959791 translocating chain-associating membrane protein
814270	-3.3246381 polymyositis/scleroderma autoantigen 1 (75kD)
884425	-3.3113482 chaperonin containing TCP1, subunit 5 (epsilon)
1874367	-3.2390912 small inducible cytokine subfamily A (Cys-Cys), member 20
742707	-3.2093502 ESTs, Weakly similar to MUC2_HUMAN MUCIN 2 PRECURSOR [H.sapiens]
280375	-3.1386611 mitochondrial ribosomal protein L15 -3.110486 PRO2000 protein -3.0906181 electron-transfer-flavoprotein, beta polypeptide
826070 624867	-3.0561472 ubiquitin specific protease 10 -3.0517117 KIAA0948 protein -2.9688171 hypothetical protein FLJ20186 -2.9361687 uncharacterized hypothalamus protein HT010
1476053	-2.9129535 RAD51 (S. cerevisiae) homolog (E coli RecA homolog)
769921	 -2.8913426 hypothetical protein MGC5585 -2.8415969 ubiquitin carrier protein E2-C -2.8216808 Homo sapiens mRNA; cDNA DKFZp762H106 (from clone DKFZp762H106)
200402 815556 1055607	-2.7995652 hypothetical protein FLJ13154 -2.7793137 hypothetical protein dJ616B8.3 -2.7693 hypothetical protein FLJ10430 -2.7538365 SUMO-1 activating enzyme subunit 1 -2.7448687 ATPase, Ca++ transporting, type 2C, member 1
770675	-2.6929924 Homo sapiens cDNA: FLJ21323 fis, clone COL02374
	-2.6909833 RAB2, member RAS oncogene family -2.6845343 ESTs, Weakly similar to S13495 pregnancy zone protein [H.sapiens]
2011138 898333	-2.6813265 KIAA0174 gene product -2.6569485 KIAA1036 protein -2.6509707 ESTs -2.6361347 brain protein I3 -2.63282 lysyl-tRNA synthetase
sd-71385	

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136722	-2.6269431 ATPase, Na+/K+ transporting, beta 3 polypeptide
951241 292936	-2.6261051 hypothetical protein FLJ10339 -2.6159344 transcription factor 4 -2.5987959 clone HQ0310 PRO0310p1 -2.5940369 hypothetical protein FLJ10468 -2.569753 budding uninhibited by benzimidazoles 1 (yeast homolog)
377368	-2.5587562 hypothetical protein FLJ10511 -2.5495557 cell death regulator aven -2.5448438 -2.539248 proteasome (prosome, macropain) subunit, beta type, 2
731223	-2.5364769 proteasome (prosome, macropain) subunit, beta type, 2
1660666 825470 1472719	
869375	-2.5249164 isocitrate dehydrogenase 2 (NADP+), mitochondrial
	-2.5097213 exostoses (multiple) 2 -2.4998247 tumor necrosis factor receptor superfamily, member 10d, decoy with truncated death domain
	 -2.488461 ribonucleotide reductase M2 polypeptide -2.4843808 methylene tetrahydrofolate dehydrogenase (NAD+ dependent), methenyltetrahydrofolate cyclohydrolase
50884	-2.4377483 dual-specificity tyrosine-(Y)-phosphorylation regulated kinase 2
1474424	-2.4345879 Homo sapiens cDNA FLJ12758 fis, clone NT2RP2001328
	-2.4324065 glycogenin -2.4311609 cyclin-dependent kinase inhibitor 3 (CDK2-associated dual specificity phosphatase)
610326 284004	-2.4310983 hypothetical protein FLJ14991 -2.4214624 -2.4185183 ESTs -2.3963227 S100 calcium-binding protein A10 (annexin II ligand, calpactin I, light polypeptide (p11))

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128711	-2.3891503 anillin (Drosophila Scraps homolog), actin binding protein
1582738	-2.3882015 uncharacterized bone marrow protein BM040
1915867	-2.3829702 Homo sapiens cDNA: FLJ21086 fis, clone CAS03272
462961	-2.3765311 ecotropic viral integration site 2A -2.3685047 dihydrofolate reductase -2.3682586 transforming, acidic coiled-coil containing protein 3
129294	-2.3605995 vanin 1 -2.3507956 ESTs -2.3342142 gamma-glutamyl hydrolase (conjugase, folylpolygammaglutamyl hydrolase) -2.333814 ESTs, Weakly similar to 2004399A chromosomal protein
815771 703707 2017415	[H.sapiens] -2.3166739 chromosome 7 open reading frame 2 -2.2865473 aspartate beta-hydroxylase -2.2832169 centromere protein A (17kD) -2.2797717 Human proteinase activated receptor-2 mRNA, 3'UTR
	-2.2761946 3'(2'), 5'-bisphosphate nucleotidase 1 -2.2753496 fucosyltransferase 4 (alpha (1,3) fucosyltransferase, myeloid-specific)
2009574	-2.2752004 Purkinje cell protein 4 -2.2646007 T brachyury (mouse) homolog -2.2563379 serine/threonine kinase 17b (apoptosis-inducing)
78869	-2.2539078 cell membrane glycoprotein, 110000M(r) (surface antigen)
	-2.2513499 sterol-C4-methyl oxidase-like -2.2489394 PDZ-binding kinase; T-cell originated protein kinase
	-2.2489064 ESTs -2.2481741 ubiquitin-conjugating enzyme E2E 2 (homologous to yeast UBC4/5)
781472	-2.2390098 Tax1 (human T-cell leukemia virus type I) binding protein 1
	-2.2375228 delta (Drosophila)-like 3 -2.2354975 minichromosome maintenance deficient (mis5, S. pombe) 6
	-2.2325695 -2.2308926 ESTs

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753320 486626 1505038	-2.2280935 epithelial V-like antigen 1 -2.2104141 hypothetical protein FLJ20533 -2.2095389 ESTs -2.2060272 hypothetical protein FLJ20171 -2.1992378 nuclear receptor co-repressor/HDAC3 complex subunit
503671	-2.1939286 Homo sapiens cDNA FLJ14368 fis, clone HEMBA1001122
712139	-2.1885269 -2.1839319 ADP-ribosylation factor-like 7 -2.1797647 integral membrane protein 3 -2.1781545 guanine nucleotide binding protein (G protein), alpha inhibiting activity polypeptide 1
183556	-2.1734918 gap junction protein, alpha 4, 37kD (connexin 37)
809557	-2.1667294 minichromosome maintenance deficient (S. cerevisiae) 3
1600239	-2.1633594 hypothetical protein MGC12992 -2.1602202 HSPC037 protein -2.1593791 olfactory receptor, family 7, subfamily E, member 47 pseudogene
839682	-2.1475459 ubiquitin-conjugating enzyme E2N (homologous to yeast UBC13)
950897	-2.1419057 Homo sapiens mRNA; cDNA DKFZp586G1922 (from clone DKFZp586G1922)
712577	-2.1411578 holocytochrome c synthase (cytochrome c heme-lyase)
241348	-2.1400285 UDP-galactose transporter related -2.1301572 prenylcysteine lyase -2.1259223 actin related protein 2/3 complex, subunit 1A (41 kD)
795498 82710 1869201 1694526	-2.1206632 lysophospholipase I -2.1190094 putative transmembrane protein -2.1184513 glutathione S-transferase A2 -2.1135906 hypothetical protein MGC2745 -2.1126121 hypothetical protein FLJ11029 -2.1072985 proteasome (prosome, macropain) 26S subunit, non-ATPase,
811590	-2.1065234 metallothionein 1H -2.1050783 hypothetical protein FLJ11100 -2.1050163 pleiomorphic adenoma gene-like 1

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	1517595 282428	-2.0965219 KIAA0175 gene product -2.0954489 ESTs, Weakly similar to A46010 X-linked retinopathy protein [H.sapiens]
	4000040	
		-2.0949091 ESTs
	746229	-2.0906355 mitogen-activated protein kinase kinase kinase kinase 4
	610326	-2.0896786 hypothetical protein MGC12992
		-2.0828168 ESTs
		-2.0796998 hypothetical protein MGC2577
		-2.0725881 extra spindle poles, S. cerevisiae, homolog of
	1410000	-2.07 2000 1 Oxtra opinialo poloco, el concincio, membro 5
	1626297	-2.0656173 hypothetical protein FLJ20509
	852829	-2.0608067 karyopherin alpha 3 (importin alpha 4)
		-2.0554449 vimentin
		-2.0519206 neurochondrin
Manager II		-2.0506628 KIAA0830 protein
	812276	-2.0458249 synuclein, alpha (non A4 component of amyloid precursor)
a		
	306841	-2.0449642 T cell receptor beta locus
	3172883	-2.0437672 ESTs, Weakly similar to 1709359A dopamine D4 receptor
		[H.sapiens]
7	624627	-2.0409259 ribonucleotide reductase M2 polypeptide
of condition of the con	486179	-2.0380168 Homo sapiens cDNA FLJ10205 fis, clone HEMBA1004954
		-2.0218598 wee1+ (S. pombe) homolog
		-2.0190035 hypothetical protein FLJ21610
		-2.0166696 myosin VB
		-2.0133834 ESTs
		-2.0067977 hypothetical protein FLJ10540
	1605426	-1.9957389 hypothetical protein FLJ13352
	810711	-1.9920785 stearoyl-CoA desaturase (delta-9-desaturase)
		4 000 4400 to the last section of the last sec
	263790	-1.9904129 isocitrate dehydrogenase 3 (NAD+) alpha
		-1.9893582 hypothetical protein PRO1068
		-1.9879196 KIAA0286 protein
	51773	-1.9875155 hypothetical protein MGC3077

Example X: Crossvalidation

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The following table shows the results of crossvalidation analysis (as described in Example I above) of various pair-wise comparisons. The two members of the pairwise comparison are indicated as "Class 1" and "Class 2" with the number of samples analyzed indicated under "N". The number of genes used to predict is indicated, along with the accuracy (in percent) for each combination. "FP" and "FN" refer to "false positive" and "false negative", respectively, as incorrectly identified for each of the two classes.

Class 1	Class 2			Genes	Accuracy	Class I		Class 2	
Name	N	Name	N			FP	FN	FP	FN
Normal	28	Abnormal	57	850	99	0	1	1	0
Normal	28	ADH	7	600	100	- 0	0	0	- 0
Normal	28	DCIS	28	1300	100	0	0	0	0
ADH	7	DCIS	28	350	97	0	1	1	0
ADH	7	DCIS	28	10	97	0	1	1	0
N+ADH	36	DCIS+IDC	49	400	98	1	1	1	1
DCIS-I	7	DCIS-III	9	300	100	0	0	0	0
IDC-I	4	IDC-III	8	300	100	0	0	0	0

References:

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All references cited herein, including patents, patent applications, and publications, are hereby incorporated by reference in their entireties, whether previously specifically incorporated or not.

Having now fully described this invention, it will be appreciated by those skilled in the art that the same can be performed within a wide range of equivalent parameters, concentrations, and conditions without departing from the spirit and scope of the invention and without undue experimentation.

While this invention has been described in connection with specific embodiments thereof, it will be understood that it is capable of further modifications. This application is intended to cover any variations, uses, or adaptations of the invention following, in general, the principles of the invention and including such departures from the present disclosure as come within known or customary practice within the art to which the invention pertains and as may be applied to the essential features hereinbefore set forth.